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57) Abstract			•
Engineered fluorescent proteins,	nucleic acids encodi	ng then	and methods of use.

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#### LONG WAVELENGTH ENGINEERED FLUORESCENT PROTEINS

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#### BACKGROUND OF THE INVENTION

This application claims the benefit of the earlier filing date of a United States provisional patent application serial number "Long Wavelength Mutant Fluorescent Proteins" and patent application serial number 08/706,408filed on August 30,1996 entitled "Long Wavelength Engineered Fluorescent Proteins" both of which are herein incorporated by reference.

This invention was made in part with Government support under grant no. MCB 9418479 awarded by the National Science Foundation. The Government may have rights in this invention.

Fluorescent molecules are attractive as reporter molecules in many assay systems because of their high sensitivity and ease of quantification. Recently, fluorescent proteins have been the focus of much attention because they can be produced in vivo by biological systems, and can be used to trace intracellular events without the need to be introduced into the cell through microinjection or permeabilization. The green fluorescent protein of Aequorea victoria is particularly interesting as a fluorescent protein. A cDNA for the protein has been cloned. (D.C. Prasher et al., "Primary structure of the Aequorea victoria green-fluorescent protein," Gene (1992) 111:229-33.) Not only can the primary amino acid sequence of the protein be expressed from the cDNA, but the expressed protein \* can fluoresce. This indicates that the protein can undergo the cyclization and oxidation believed to be necessary for fluorescence. Aeguorea green fluorescent protein ("GFP") is a stable, proteolysis-resistant single chain of 238 residues and has two absorption maxima at around 395 and 475 nm. The relative amplitudes of these two peaks is sensitive to environmental factors (W. W. Ward, Bioluminescence and Chemiluminescence (M. A. DeLuca and W. D. McElroy, eds) Academic Press pp. 235-242 (1981); W. W. Ward & S. H. Bokman Biochemistry 21:4535-4540 (1982); W. W. Ward et al. Photochem. Photobiol. 35:803-808 (1982)) and illumination history (A. B. Cubitt et al. Trends Biochem, Sci. 20:448-455 (1995)), presumably reflecting two or more ground states. Excitation at the primary absorption peak of 395 nm yields an emission maximum at 508 nm with a quantum yield of 0.72-0.85 (O. Shimomura and F.H. Johnson J. Cell. Comp. Physiol. 59:223 (1962);

J. G. Morin and J. W. Hastings, J. Cell. Physiol. 77:313 (1971); H. Morise et al. Biochemistry 13:2656 (1974); W. W. Ward Photochem. Photobiol. Reviews (Smith, K. C. ed.) 4:1 (1979); A. B. Cubitt et al. Trends Biochem. Sci. 20:448-455 (1995); D. C. Prasher Trends Genet. 11:320-323 (1995); M. Challie Photochem. Photobiol. 62:651-656 (1995); W. W. Ward. Bioluminescence and Chemilluminescence (M. A. DeLuca and W. D. McElroy, eds) Academic Press pp. 235-242 (1981); W. W. Ward & S. H. Bokman Biochemistry 21:4535-4540 (1982); W. W. Ward et al. Photochem. Photobiol. 35:803-808 (1982)). The fluorophore results from the autocatalytic cyclization of the polypeptide backbone between residues Ser<sup>55</sup> and GiJ<sup>57</sup> and oxidation of the □-8 bond of Tyr<sup>66</sup> (A. B. Cubitt et al. Trends Biochem. Sci. 20:448-4455 (1995); C. W. Cody et al. Biochemistry

32:1212-1218 (1993); R. Heim et al. Proc. Natl. Acad. Sci. USA 91:12501-12504 (1994)). Mutation of Ser<sup>66</sup> to Thr (S65T) simplifies the excitation spectrum to a single peak at 488 nm of enhanced amplitude (R. Heim et al. Nature 373:664-665 (1995)), which no longer gives signs of conformational isomers (A. B. Cubitt et al. Trends Biochem. Sci. 20:448-455 (1995)).

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Fluorescent proteins have been used as markers of gene expression, tracers of cell lineage and as fusion tags to monitor protein localization within living cells. (M. Chalfie et al., "Green fluorescent protein as a marker for gene expression," Science 263:802-805; A.B. Cubitt et al., "Understanding, improving and using green fluorescent proteins,"

20 TIBS 20, November 1995, pp. 448-455. U.S. patent 5,491,084, M. Chalfie and D. Prasher. Furthermore, engineered versions of Aequorea green fluorescent protein have been identified that exhibit altered fluorescence characteristics, including altered excitation and emission maxima, as well as excitation and emission spectra of different shapes. (R. Heim et al., "Wavelength mutations and posttranslational autoxidation of green fluorescent protein," Proc. Natl. Acad. Sci. USA, (1994) 91:12501-04; R. Heim et al., "Improved green fluorescence," Nature (1995) 373:663-665.) These properties add variety and utility to the arsenal of biologically based fluorescent indicators.

There is a need for engineered fluorescent proteins with varied fluorescent properties.

#### BRIEF DESCRIPTION OF THE DRAWINGS

Figs. 1A-1B. (A) Schematic drawing of the backbone of GFP produced by

Molscript (I.P. Kraulis, J. Appl. Cryst., 24:946 (1991)). The chromophore is shown as a ball and stick model. (B) Schematic drawing of the overall fold of GFP. Approximate residue numbers mark the beginning and ending of the secondary structure elements.

Figs. 2A-2C. (A) Stereo drawing of the chromophore and residues in the immediate vicinity. Carbon atoms are drawn as open circles, oxygen is filled and nitrogen is shaded. Solvent molecules are shown as isolated filled circles. (B) Portion of the final  $2F_s$ -felectron density map contoured at  $1.0\,\Box$ , showing the electron density surrounding the chromophore. (C) Schematic diagram showing the first and second spheres of coordination of the chromophore. Hydrogen bonds are shown as dashed lines and have the indicated lengths in  $\dot{A}$ . Inset: proposed structure of the carbinolamine intermediate that is presumably formed during generation of the chromophore.

Fig. 3 depicts the nucleotide sequence (SEQ ID NO:1) and deduced amino acid sequence (SEQ ID NO:2) of an Aequorea green fluorescent protein.

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Fig. 4 depicts the nucleotide sequence (SEQ ID NO:3) and deduced amino acid sequence (SEQ ID NO:4) of the engineered *Aequorea*-related fluorescent protein S65G/S72A/T203Y utilizing preferred mammalian codons and optimal Kozak sequence.

Figs. 5-1 to 5-28 present the coordinates for the crystal structure of Aeguorea-related green fluorescent protein S65T.

Fig. 6 shows the fluorescence excitation and emission spectra for engineered fluorescent proteins 20A and 10C (Table F). The vertical line at 528 nm compares the emission maxima of 10C, to the left of the line, and 20A, to the right of the line.

#### SUMMARY OF THE INVENTION

This invention provides functional engineered fluorescent proteins with

varied fluorescence characteristics that can be easily distinguished from currently existing
green and blue fluorescent proteins. Such engineered fluorescent proteins enable the
simultaneous measurement of two or more processes within cells and can be used as
fluorescence energy donors or acceptors when used to monitor protein-protein interactions
through FRET. Longer wavelength engineered fluorescent proteins are particularly useful

because photodynamic toxicity and auto-fluorescence of cells are significantly reduced at
longer wavelengths. In particular, the introduction of the substitution T203X, wherein X is
an aromatic amino acid, results in an increase in the excitation and emission wavelength

maxima of Aequorea-related fluorescent proteins.

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In one aspect, this invention provides a nucleic acid molecule comprising a nucleotide sequence encoding a functional engineered fluorescent protein whose amino acid sequence is substantially identical to the amino acid sequence of Aequorea green fluorescent protein (SEQ ID NO:2) and which differs from SEQ ID NO:2 by at least an amino acid substitution located no more than about 0.5 nm from the chromophore of the engineered fluorescent protein, wherein the substitution alters the electronic environment of the chromophore, whereby the functional engineered fluorescent protein has a different fluorescent property than Aeguorea green fluorescent protein.

In one aspect this invention provides a nucleic acid molecule comprising a nucleotide sequence encoding a functional engineered fluorescent protein whose amino acid sequence is substantially identical to the amino acid sequence of Aequorea green fluorescent protein (SEQ ID NO:2) and which differs from SEQ ID NO:2 by at least a substitution at T203 and, in particular, T203X, wherein X is an aromatic amino acid selected from H, Y, W or F, said functional engineered fluorescent protein having a different fluorescent property than Aequorea green fluorescent protein. In one embodiment, the amino acid sequence further comprises a substitution at S65, wherein the substitution is selected from S65G, S65T, S65A, S65L, S65C, S65V and S65I. In another embodiment, the amino acid sequence differs by no more than the substitutions S65T/T203H; S65T/T203Y; S72A/F64L/S65G/T203Y; S65G/V68L/Q69K/S72A/T203Y; S72A/S65G/V68L/T203Y; S65G/S72A/T203Y; or S65G/S72A/T203W. In another embodiment, the amino acid sequence further comprises a substitution at Y66, wherein the substitution is selected from Y66H, Y66F, and Y66W. In another embodiment, the amino acid sequence further comprises a mutation from Table A. In another embodiment, the amino acid sequence further comprises a folding mutation. In another embodiment, the nucleotide sequence encoding the protein differs from the nucleotide sequence of SEO ID NO:1 by the substitution of at least one codon by a preferred mammalian codon. In another embodiment, the nucleic acid molecule encodes a fusion protein wherein the fusion protein comprises a

In another aspect, this invention provides a nucleic acid molecule comprising a nucleotide sequence encoding a functional engineered fluorescent protein whose amino acid sequence is substantially identical to the amino acid sequence of *Aequorea* green

polypeptide of interest and the functional engineered fluorescent protein.

fluorescent protein (SEQ ID NO:2) and which differs from SEQ ID NO:2 by at least an amino acid substitution at L42, V61, T62, V68, Q69, Q94, N121, Y145, H148, V150, F165, 1167, O183, N185, L220, E222 (not E222G), or V224, said functional engineered fluorescent protein having a different fluorescent property than Aequorea green fluorescent protein. In one embodiment, amino acid substitution is:

L42X, wherein X is selected from C, F, H, W and Y.

V61X, wherein X is selected from F. Y. H and C.

T62X, wherein X is selected from A. V. F. S. D. N. O. Y. H and C.

V68X, wherein X is selected from F, Y and H,

O69X, wherein X is selected from K, R, E and G, 10

O94X, wherein X is selected from D, E, H, K and N.

N121X, wherein X is selected from F. H. W and Y.

Y145X, wherein X is selected from W, C, F, L, E, H, K and O.

H148X, wherein X is selected from F, Y, N, K, O and R,

V150X, wherein X is selected from F, Y and H. 15

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F165X, wherein X is selected from H, Q, W and Y,

I167X, wherein X is selected from F, Y and H.

O183X, wherein X is selected from H, Y, E and K,

N185X, wherein X is selected from D, E, H, K and O,

L220X, wherein X is selected from H, N, O and T.

E222X, wherein X is selected from N and Q, or

V224X, wherein X is selected from H. N. O. T. F. W and Y.

In a further aspect, this invention provides an expression vector comprising 25 expression control sequences operatively linked to any of the aforementioned nucleic acid molecules. In a further aspect, this invention provides a recombinant host cell comprising the aforementioned expression vector.

In another aspect, this invention provides a functional engineered fluorescent protein whose amino acid sequence is substantially identical to the amino acid sequence of Aeguorea green fluorescent protein (SEQ ID NO:2) and which differs from SEO ID NO:2 by at least an amino acid substitution located no more than about 0.5 nm from the chromophore of the engineered fluorescent protein, wherein the substitution alters the

electronic environment of the chromophore, whereby the functional engineered fluorescent protein has a different fluorescent property than Aequorea green fluorescent protein.

In another aspect, this invention provides a functional engineered fluorescent protein whose amino acid sequence is substantially identical to the amino acid sequence of Aequorea green fluorescent protein (SEQ ID NO:2) and which differs from SEQ ID NO:2 by at least the amino acid substitution at T203, and in particular, T203X, wherein X is an aromatic amino acid selected from H. Y. W or F. said functional engineered fluorescent protein having a different fluorescent property than Aequorea green fluorescent protein. In one embodiment, the amino acid sequence further comprises a substitution at S65, wherein the substitution is selected from S65G, S65T, S65A, S65L, S65C, S65V and S65I. In another embodiment, the amino acid sequence differs by no more than the substitutions S65T/T203H; S65T/T203Y; S72A/F64L/S65G/T203Y; S72A/S65G/V68L/T203Y; S65G/V68L/O69K/S72A/T203Y: S65G/S72A/T203Y: or S65G/S72A/T203W. In another embodiment, the amino acid sequence further comprises a substitution at Y66, wherein the substitution is selected from Y66H, Y66F, and Y66W. In another embodiment, the amino acid sequence further comprises a folding mutation. In another embodiment, the engineered fluorescent protein is part of a fusion protein wherein the fusion protein comprises a polypeptide of interest and the functional engineered fluorescent protein.

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In another aspect this invention provides a functional engineered fluorescent protein whose amino acid sequence is substantially identical to the amino acid sequence of Aequorea green fluorescent protein (SEQ ID NO:2) and which differs from SEQ ID NO:2 by at least an amino acid substitution at L42, V61, T62, V68, Q69, Q94, N121, Y145, H148, V150, F165, I167, Q183, N185, L220, E222, or V224, said functional engineered fluorescent protein having a different fluorescent property than Aequorea green fluorescent protein.

In another aspect, this invention provides a fluorescently labelled antibody comprising an antibody coupled to any of the aforementioned functional engineered fluorescent proteins. In one embodiment, the fluorescently labelled antibody is a fusion protein wherein the fusion protein comprises the antibody fused to the functional engineered fluorescent protein.

In another aspect, this invention provides a nucleic acid molecule comprising a nucleotide sequence encoding an antibody fused to a nucleotide sequence encoding a 1.0

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functional engineered fluorescent protein of this invention.

In another aspect, this invention provides a fluorescently labelled nucleic acid probe comprising a nucleic acid probe coupled to a functional engineered fluorescent protein whose amino acid sequence of this invention. The fusion can be through a linker peptide.

In another aspect, this invention provides a method for determining whether a mixture contains a target comprising contacting the mixture with a fluorescently labelled probe comprising a probe and a functional engineered fluorescent protein of this invention; and determining whether the target has bound to the probe. In one embodiment, the target molecule is captured on a solid matrix.

In another aspect, this invention provides a method for engineering a functional engineered fluorescent protein having a fluorescent property different than Aequorea green fluorescent protein, comprising substituting an amino acid that is located no more than 0.5 nm from any atom in the chromophore of an Aequorea-related green fluorescent protein with another amino acid; whereby the substitution alters a fluorescent property of the protein. In one embodiment, the amino acid substitution alters the electronic environment of the chromophore.

In another aspect, this invention provides a method for engineering a functional engineered fluorescent protein having a different fluorescent property than Aequorea green fluorescent protein comprising substituting amino acids in a loop domain of an Aequorea-related green fluorescent protein with amino acids so as to create a consensus\* sequence for phosphorylation or for proteolysis.

In another aspect, this invention provides a method for producing fluorescence resonance energy transfer comprising providing a donor molecule comprising a functional engineered fluorescent protein this invention; providing an appropriate acceptor molecule for the fluorescent protein; and bringing the donor molecule and the acceptor molecule into sufficiently close contact to allow fluorescence resonance energy transfer.

In another aspect, this invention provides a method for producing fluorescence resonance energy transfer comprising providing an acceptor molecule comprising a functional engineered fluorescent protein of this invention; providing an appropriate donor molecule for the fluorescent protein; and bringing the donor molecule and the acceptor molecule into sufficiently close contact to allow fluorescence resonance energy

transfer. In one embodiment, the donor molecule is a engineered fluorescent protein whose amino acid sequence comprises the substitution T203I and the acceptor molecule is an engineered fluorescent protein whose amino acid sequence comprises the substitution T203X, wherein X is an aromatic amino acid selected from H, Y, W or F, said functional engineered fluorescent protein having a different fluorescent property than Aequorea green fluorescent protein.

In another aspect, this invention provides a crystal of a protein comprising a fluorescent protein with an amino acid sequence substantially identical to SEQ ID NO: 2, wherein said crystal diffracts with at least a 2.0 to 3.0 angstrom resolution.

In another embodiment, this invention provides computational method of designing a fluoresent protein comprising determining from a three dimensional model of a crystallized fluorescent protein comprising a fluorescent protein with a bound ligand, at least one interacting amino acid of the fluorescent protein that interacts with at least one first chemical moiety of the ligand, and selecting at least one chemical modification of the first chemical moiety to produce a second chemical moiety with a structure to either decrease or increase an interaction between the interacting amino acid and the second chemical moiety compared to the interaction between the interacting amino acid and the first chemical moiety.

In another embediment, this invention provides a computational method of modeling the three dimensional structure of a fluorescent protein comprising determining a three dimensional relationship between at least two atoms listed in the atomic coordinates of\* Figs. 5-1 to 5-28.

In another embodiment, this invention provides a device comprising a storage device and, stored in the device, at least 10 atomic coordinates selected from the atomic coordinates listed in Figs. 5-1 to 5-28. In one embodiment, the storage device is a computer readable device that stores code that receives as input the atomic coordinates. In another embodiment, the computer readable device is a floppy disk or a hard drive.

#### DETAILED DESCRIPTION OF THE INVENTION

#### 30 I. DEFINITIONS

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Unless defined otherwise, all technical and scientific terms used herein have the same meaning as commonly understood by those of ordinary skill in the art to which WO 98/06737 PCT/US97/14593

this invention belongs. Although any methods and materials similar or equivalent to those described herein can be used in the practice or testing of the present invention, the preferred methods and materials are described. For purposes of the present invention, the following terms are defined below.

"Binding pair" refers to two moieties (e.g. chemical or biochemical) that have an affinity for one another. Examples of binding pairs include antigen/antibodies, lectin/avidin, target polynucleotide/probe oligonucleotide, antibody/anti-antibody, receptor/ligand, enzyme/ligand and the like. "One member of a binding pair" refers to one moiety of the pair, such as an antigen or ligand.

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"Nucleic acid" refers to a deoxyribonucleotide or ribonucleotide polymer in either single- or double-stranded form, and, unless otherwise limited, encompasses known analogs of natural nucleotides that can function in a similar manner as naturally occurring nucleotides. It will be understood that when a nucleic acid molecule is represented by a DNA sequence, this also includes RNA molecules having the corresponding RNA sequence in which "U" replaces "T."

"Recombinant nucleic acid molecule" refers to a nucleic acid molecule which is not naturally occurring, and which comprises two nucleotide sequences which are not naturally joined together. Recombinant nucleic acid molecules are produced by artificial recombination, e.g., genetic engineering techniques or chemical synthesis.

Reference to a nucleotide sequence "encoding" a polypeptide means that the sequence, upon transcription and translation of mRNA, produces the polypeptide. This includes both the coding strand, whose nucleotide sequence is identical to mRNA and whose sequence is usually provided in the sequence listing, as well as its complementary strand, which is used as the template for transcription. As any person skilled in the art recognizes, this also includes all degenerate nucleotide sequences encoding the same amino acid sequence. Nucleotide sequences encoding a polypeptide include sequences containing introns.

"Expression control sequences" refers to nucleotide sequences that regulate the expression of a nucleotide sequence to which they are operatively linked. Expression control sequences are "operatively linked" to a nucleotide sequence when the expression control sequences control and regulate the transcription and, as appropriate, translation of the nucleotide sequence. Thus, expression control sequences can include appropriate

promoters, enhancers, transcription terminators, a start codon (i.e., ATG) in front of a protein-encoding gene, splicing signals for introns, maintenance of the correct reading frame of that gene to permit proper translation of the mRNA, and stop codons.

"Naturally-occurring" as used herein, as applied to an object, refers to the fact that an object can be found in nature. For example, a polypeptide or polynucleotide sequence that is present in an organism (including viruses) that can be isolated from a source in nature and which has not been intentionally modified by man in the laboratory is naturally-occurring.

"Operably linked" refers to a juxtaposition wherein the components so described are in a relationship permitting them to function in their intended manner. A control sequence "operably linked" to a coding sequence is ligated in such a way that expression of the coding sequence is achieved under conditions compatible with the control sequences, such as when the appropriate molecules (e.g., inducers and polymerases) are bound to the control or regulatory sequence(s).

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"Control sequence" refers to polynucleotide sequences which are necessary to effect the expression of coding and non-coding sequences to which they are ligated. The nature of such control sequences differs depending upon the host organism; in prokaryotes, such control sequences generally include promoter, ribosomal binding site, and transcription termination sequence; in sukaryotes, generally, such control sequences include promoters and transcription termination sequence. The term "control sequences" is intended to include, at a minimum, components whose presence can influence expression, and can also include additional components whose presence is advantageous, for example, leader sequences and fusion partner sequences.

"Isolated polynucleotide" refers a polynucleotide of genomic, cDNA, or synthetic origin or some combination there of, which by virtue of its origin the "isolated polynucleotide" (1) is not associated with the cell in which the "isolated polynucleotide" is found in nature, or (2) is operably linked to a polynucleotide which it is not linked to in nature.

"Polynucleotide" refers to a polymeric form of nucleotides of at least 10 bases in length, either ribonucleotides or deoxynucleotides or a modified form of either type of nucleotide. The term includes single and double stranded forms of DNA.

The term "probe" refers to a substance that specifically binds to another substance (a "target"). Probes include, for example, antibodies, nucleic acids, receptors and

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"Modulation" refers to the capacity to either enhance or inhibit a functional property of biological activity or process (e.g., enzyme activity or receptor binding); such enhancement or inhibition may be contingent on the occurrence of a specific event, such as activation of a signal transduction pathway, and/or may be manifest only in particular cell types.

The term "modulator" refers to a chemical (naturally occurring or non-naturally occurring), such as a synthetic molecule (e.g., nucleic acid, protein, non-peptide, or organic molecule), or an extract made from biological materials such as bacteria, plants, fungi, or animal (particularly mammalian) cells or tissues. Modulators can be evaluated for potential activity as inhibitors or activators (directly or indirectly) of a biological process or processes (e.g., agonist, partial antagonist, partial agonist, inverse agonist, antagonist, antineoplastic agents, cytotoxic agents, inhibitors of neoplastic transformation or cell proliferation, cell proliferation-promoting agents, and the like) by inclusion in screening assays described herein. The activity of a modulator may be known, unknown or partially known.

The term "test chemical" refers to a chemical to be tested by one or more screening method(s) of the invention as a putative modulator. A test chemical is usually not known to bind to the target of interest. The term "control test chemical" refers to a chemical known to bind to the target (e.g., a known agonist, antagonist, partial agonist or inverse agonist).

Usually, various predetermined concentrations of test chemicals are used for screening, such as .01 µM, .1 µM, 1.0 µM, and 10.0 µM.

The term "target" refers to a biochemical entity involved a biological process.

Targets are typically proteins that play a useful role in the physiology or biology of an organism. A therapeutic chemical binds to target to alter or modulate its function. As used herein targets can include cell surface receptors, G-proteins, kinases, ion channels, phopholipases and other proteins mentioned herein.

The term "label" refers to a composition detectable by spectroscopic, photochemical, biochemical, immunochemical, or chemical means. For example, useful labels include <sup>32</sup>P, fluorescent dyes, fluorescent proteins, electron-dense reagents, enzymes (e.g., as commonly used in an ELISA), biotin, dioxigenin, or haptens and proteins for which antisera or monoclonal antibodies are available. For example, polypeptides of this invention can be made as detectible labels, by e.g., incorporating a them as into a polypeptide, and

used to label antibodies specifically reactive with the polypeptide. A label often generates a measurable signal, such as radioactivity, fluorescent light or enzyme activity, which can be used to quantitate the amount of bound label.

The term "nucleic acid probe" refers to a nucleic acid molecule that binds to a specific sequence or sub-sequence of another nucleic acid molecule. A probe is preferably a nucleic acid molecule that binds through complementary base pairing to the full sequence or to a sub-sequence of a target nucleic acid. It will be understood that probes may bind target sequences lacking complete complementarity with the probe sequence depending upon the stringency of the hybridization conditions. Probes are preferably directly labelled as with isotopes, chromophores, lumiphores, chromogens, fluorescent proteins, or indirectly labelled such as with biotin to which a streptavidin complex may later bind. By assaying for the presence or absence of the probe, one can detect the presence or absence of the select sequence or sub-sequence.

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A "labeled nucleic acid probe" is a nucleic acid probe that is bound, either covalently, through a linker, or through ionic, van der Waals or hydrogen bonds to a label such that the presence of the probe may be detected by detecting the presence of the label bound to the probe.

The terms "polypeptide" and "protein" refers to a polymer of amino acid residues. The terms apply to amino acid polymers in which one or more amino acid residue is an artificial chemical analogue of a corresponding naturally occurring amino acid, as well as to naturally occurring amino acid polymers. The term "recombinant protein" refers to a protein that is produced by expression of a nucleotide sequence encoding the amino acid sequence of the protein from a recombinant DNA molecule.

The term "recombinant host cell" refers to a cell that comprises a recombinant nucleic acid molecule. Thus, for example, recombinant host cells can express genes that are not found within the native (non-recombinant) form of the cell.

The terms "isolated" "purified" or "biologically pure" refer to material which is substantially or essentially free from components which normally accompany it as found in its native state. Purity and homogeneity are typically determined using analytical chemistry techniques such as polyacrylamide gel electrophoresis or high performance liquid chromatography. A protein or nucleic acid molecule which is the predominant protein or nucleic acid species present in a preparation is substantially purified. Generally, an isolated

protein or nucleic acid molecule will comprise more than 80% of all macromolecular species present in the preparation. Preferably, the protein is purified to represent greater than 90% of all macromolecular species present. More preferably the protein is purified to greater than 95%, and most preferably the protein is purified to essential homogeneity, wherein other macromolecular species are not detected by conventional techniques.

The term "naturally-occurring" as applied to an object refers to the fact that an object can be found in nature. For example, a polypeptide or polynucleotide sequence that is present in an organism (including viruses) that can be isolated from a source in nature and which has not been intentionally modified by man in the laboratory is naturally-occurring.

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The term "antibody" refers to a polypeptide substantially encoded by an immunoglobulin gene or immunoglobulin genes, or fragments thereof, which specifically bind and recognize an analyte (antigen). The recognized immunoglobulin genes include the kappa, lambda, alpha, gamma, delta, epsilon and mu constant region genes, as well as the myriad immunoglobulin variable region genes. Antibodies exist, e.g., as intact immunoglobulins or as a number of well characterized fragments produced by digestion with various peptidases. This includes, e.g., Fab' and F(ab)'<sub>2</sub> fragments. The term "antibody," as used herein, also includes antibody fragments either produced by the modification of whole antibodies or those synthesized de novo using recombinant DNA methodologies.

The term "immunoassay" refers to an assay that utilizes an antibody to specifically bind an analyte. The immunoassay is characterized by the use of specific binding properties of a particular antibody to isolate, target, and/or quantify the analyte.

The term "identical" in the context of two nucleic acid or polypeptide sequences refers to the residues in the two sequences which are the same when aligned for maximum correspondence. When percentage of sequence identity is used in reference to proteins or peptides it is recognized that residue positions which are not identical often differ by conservative amino acid substitutions, where amino acids residues are substituted for other amino acid residues with similar chemical properties (e.g. charge or hydrophobicity) and therefore do not change the functional properties of the molecule. Where sequences differ in conservative substitutions, the percent sequence identity may be adjusted upwards to correct for the conservative nature of the substitution. Means for

making this adjustment are well known to those of skill in the art. Typically this involves scoring a conservative substitution as a partial rather than a full mismatch, thereby increasing the percentage sequence identity. Thus, for example, where an identical amino acid is given a score of 1 and a non-conservative substitution is given a score of zero, a conservative substitution is given a score between zero and 1. The scoring of conservative substitutions is calculated, e.g., according to known algorithm. See, e.g., Meyers and Miller, Computer Applic. Biol. Sci., 4: 11-17 (1988); Smith and Waterman (1981) Adv. Appl. Math. 2: 482; Needleman and Wunsch (1970) J. Mol. Biol. 48: 443; Pearson and Lipman (1988) Proc. Natl. Acad. Sci. USA 85: 2444; Higgins and Sharp (1988) Gene, 73: 237-244 and Higgins and Sharp (1989) CABIOS 5: 151-153; Corpet, et al. (1988) Nucleic Acids Research 16, 10881-90; Huang, et al. (1992) Computer Applications in the Biosciences 8, 155-65, and Pearson, et al. (1994) Methods in Molecular Biology 24, 307-31. Alignment is also often performed by inspection and manual alignment.

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"Conservatively modified variations" of a particular nucleic acid sequence refers to those nucleic acids which encode identical or essentially identical amino acid sequences, or where the nucleic acid does not encode an amino acid sequence, to essentially identical sequences. Because of the degeneracy of the genetic code, a large number of functionally identical nucleic acids encode any given polypeptide. For instance, the codons CGU, CGC, CGA, CGG, AGA, and AGG all encode the amino acid arginine. Thus, at every position where an arginine is specified by a codon, the codon can be altered to any of the corresponding codons described without altering the encoded polypeptide. Such nucleic \* acid variations are "silent variations," which are one species of "conservatively modified variations." Every nucleic acid sequence herein which encodes a polypeptide also describes every possible silent variation. One of skill will recognize that each codon in a nucleic acid (except AUG, which is ordinarily the only codon for methionine) can be modified to yield a functionally identical molecule by standard techniques. Accordingly, each "silent variation" of a nucleic acid which encodes a polypeptide is implicit in each described sequence. Furthermore, one of skill will recognize that individual substitutions, deletions or additions which alter, add or delete a single amino acid or a small percentage of amino acids (typically less than 5%, more typically less than 1%) in an encoded sequence are "conservatively modified variations" where the alterations result in the substitution of an amino acid with a chemically similar amino acid. Conservative amino acid substitutions

providing functionally similar amino acids are well known in the art. The following six groups each contain amino acids that are conservative substitutions for one another:

- 1) Alanine (A), Serine (S), Threonine (T);
- 2) Aspartic acid (D), Glutamic acid (E);
- 3) Asparagine (N), Glutamine (Q);
  - 4) Arginine (R), Lysine (K);

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- 5) Isoleucine (I), Leucine (L), Methionine (M), Valine (V); and
- 6) Phenylalanine (F), Tyrosine (Y), Tryptophan (W).

The term "complementary" means that one nucleic acid molecule has the sequence of the binding partner of another nucleic acid molecule. Thus, the sequence 5'-ATGC-3' is complementary to the sequence 5'-GCAT-3'.

An amino acid sequence or a nucleotide sequence is "substantially identical" or "substantially similar" to a reference sequence if the amino acid sequence or nucleotide sequence has at least 80% sequence identity with the reference sequence over a given comparison window. Thus, substantially similar sequences include those having, for example, at least 85% sequence identity, at least 90% sequence identity, at least 95% sequence identity or at least 99% sequence identity. Two sequences that are identical to each other are, of course, also substantially identical.

A subject nucleotide sequence is "substantially complementary" to a reference nucleotide sequence if the complement of the subject nucleotide sequence is substantially identical to the reference nucleotide sequence.

The term "stringent conditions" refers to a temperature and ionic conditions used in nucleic acid hybridization. Stringent conditions are sequence dependent and are different under different environmental parameters. Generally, stringent conditions are selected to be about  $5\Box C$  to  $20\Box C$  lower than the thermal melting point  $(T_m)$  for the specific sequence at a defined ionic strength and pH. The  $T_m$  is the temperature (under defined ionic strength and pH) at which 50% of the target sequence hybridizes to a perfectly matched probe.

The term "allelic variants" refers to polymorphic forms of a gene at a particular genetic locus, as well as cDNAs derived from mRNA transcripts of the genes and the polypeptides encoded by them.

The term "preferred mammalian codon" refers to the subset of codons from

among the set of codons encoding an amino acid that are most frequently used in proteins expressed in mammalian cells as chosen from the following list:

Amino Acid Preferred codons for high level mammalian expression

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	Gly	GGC,GGG
	Glu	GAG
	Asp	GAC
	Val	GUG,GUC
10	Ala	GCC,GCU
	Ser	AGC,UCC
	Lys	AAG
	Asn	AAC
	Met	AUG
15	Ile	AUC
	Thr	ACC
	Trp	UGG
	Cys	UGC
	Тут	UAU,UAC
20	Leu	CUG
	Phe	UUC
	Arg	CGC,AGG,AGA
	Gln	CAG
	His	CAC
25	Pro	CCC

Fluorescent molecules are useful in fluorescence resonance energy transfer ("FRET"). FRET involves a donor molecule and an acceptor molecule. To optimize the efficiency and detectability of FRET between a donor and acceptor molecule, several factors need to be balanced. The emission spectrum of the donor should overlap as much as possible with the excitation spectrum of the acceptor to maximize the overlap integral. Also, the quantum yield of the donor moiety and the extinction coefficient of the acceptor should likewise be as high as possible to maximize R<sub>0</sub>, the distance at which energy transfer efficiency is 50%. However, the excitation spectra of the donor and acceptor should overlap as little as possible so that a wavelength region can be found at which the donor can be excited efficiently without directly exciting the acceptor. Fluorescence arising from firect excitation of the acceptor is difficult to distinguish from fluorescence arising from FRET. Similarly, the emission spectra of the donor and acceptor should overlap as little as possible so that the two emissions can be clearly distinguished. High fluorescence quantum yield of

the acceptor mojety is desirable if the emission from the acceptor is to be measured either as the sole readout or as part of an emission ratio. One factor to be considered in choosing the donor and acceptor pair is the efficiency of fluorescence resonance energy transfer between them. Preferably, the efficiency of FRET between the donor and acceptor is at least 10%. more preferably at least 50% and even more preferably at least 80%.

The term "fluorescent property" refers to the molar extinction coefficient at an appropriate excitation wavelength, the fluorescence quantum efficiency, the shape of the excitation spectrum or emission spectrum, the excitation wavelength maximum and emission wavelength maximum, the ratio of excitation amplitudes at two different wavelengths, the ratio of emission amplitudes at two different wavelengths, the excited state 10 lifetime, or the fluorescence anisotropy. A measurable difference in any one of these properties between wild-type Aequorea GFP and the mutant form is useful. A measurable difference can be determined by determining the amount of any quantitative fluorescent property, e.g., the amount of fluorescence at a particular wavelength, or the integral of 15 fluorescence over the emission spectrum. Determining ratios of excitation amplitude or emission amplitude at two different wavelengths ("excitation amplitude ratioing" and "emission amplitude ratioing", respectively) are particularly advantageous because the ratioing process provides an internal reference and cancels out variations in the absolute brightness of the excitation source, the sensitivity of the detector, and light scattering or quenching by the sample.

#### LONG WAVELENGTH ENGINEERED FLUORESCENT PROTEINS H

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proteins using the same standards.

#### A. Fluorescent Proteins

As used herein, the term "fluorescent protein" refers to any protein capable of fluorescence when excited with appropriate electromagnetic radiation. This includes fluorescent proteins whose amino acid sequences are either naturally occurring or engineered (i.e., analogs or mutants). Many cnidarians use green fluorescent proteins ("GFPs") as energy-transfer acceptors in bioluminescence. A "green fluorescent protein," as used herein, is a protein that fluoresces green light. Similarly, "blue fluorescent proteins" fluoresce blue light and "red fluorescent proteins" fluoresce red light. GFPs have been isolated from the Pacific Northwest jellyfish, Aequorea victoria, the sea pansy, Renilla reniformis, and Phialidium gregarium. W.W. Ward et al., Photochem. Photobiol., 35:803-808 (1982); L.D. Levine et al., Comp. Biochem. Physiol., 72B:77-85 (1982)

A variety of Aequorea-related fluorescent proteins having useful excitation and emission spectra have been engineered by modifying the amino acid sequence of a naturally occurring GFP from Aequorea victoria. (D.C. Prasher et al., Gene, 111:229-233 (1992); R. Heim et al., Proc. Natl. Acad. Sci., USA, 91:12501-04 (1994); U.S. patent application 08/337,915, filed November 10, 1994; International application PCTA:S95/14692, filed 11/10/95.)

As used herein, a fluorescent protein is an "Aequorea-related fluorescent protein" if any contiguous sequence of 150 amino acids of the fluorescent protein has at least 85% sequence identity with an amino acid sequence, either contiguous or non-contiguous, from the 238 amino-acid wild-type Aequorea green fluorescent protein of Fig. 3 • (SEQ ID NO:2). More preferably, a fluorescent protein is an Aequorea-related fluorescent protein if any contiguous sequence of 200 amino acids of the fluorescent protein has at least 95% sequence identity with an amino acid sequence, either contiguous or non-contiguous, from the wild type Aequorea green fluorescent protein of Fig. 3 (SEQ ID NO:2). Similarly,

the fluorescent protein may be related to Renilla or Phialidium wild-type fluorescent

Aequorea-related fluorescent proteins include, for example and without limitation, wild-type (native) Aequorea victoria GFP (D.C. Prasher et al., "Primary structure of the Aequorea victoria green fluorescent protein," Gene, (1992) 111:229-33), whose nucleotide sequence (SEQ ID NO:1) and deduced amino acid sequence (SEQ ID NO:2) are presented in Fig. 3; allelic variants of this sequence, e.g., O80R, which has the glutamine

residue at position 80 substituted with arginine (M. Chalfie et al., Science, (1994) 263:802-805); those engineered Aequorea-related fluorescent proteins described herein, e.g., in Table A or Table F, variants that include one or more folding mutations and fragments of these proteins that are fluorescent, such as Aequorea green fluorescent protein from which the two amino-terminal amino acids have been removed. Several of these contain different aromatic amino acids within the central chromophore and fluoresce at a distinctly shorter wavelength than wild type species. For example, engineered proteins P4 and P4-3 contain (in addition to other mutations) the substitution Y66H, whereas W2 and W7 contain (in addition to other mutations) Y66W. Other mutations both close to the chromophore region of the protein and remote from it in primary sequence may affect the spectral properties of GFP and are listed in the first part of the table below.

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TABLE A

Clone	Mutation(s)	Excitation max (nm)	Emission max (nm)	Extinct. Coeff. (M-1cm-1)	Quantum yield
Wild type	None	395 (475)	508	21,000 (7,150)	0.77
P4	Y66H	383	447	13,500	0.21
P4-3	Y66H Y145F	381	445	14,000	0.38
W7	Y66W N146I M153T V163A N212K	433 (453)	475 (501)	18,000 (17,100)	0.67
W2	Y66W I123V Y145H H148R M153T V163A N212K	432 (453)	480	10,000 (9,600)	0.72
S65T	S65T	489	511	39,200	0.68
P4-1	S65T M153A	504 (396)	514	14,500 (8,600)	0.53

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	K238E			
S65A	S65A	471	504	
S65C	S65C	479	507	
S65L	S65L	484	510	
Y66F	Y66F	360	442	
Y66W	Y66W	458	480	

Additional mutations in Aequorea-related fluorescent proteins, referred to as "folding mutations," improve the ability of fluorescent proteins to fold at higher temperatures, and to be more fluorescent when expressed in mammalian cells, but have little or no effect on the peak wavelengths of excitation and emission. It should be noted that these may be combined with mutations that influence the spectral properties of GFP to produce proteins with altered spectral and folding properties. Folding mutations include: F64L, V68L, S72A, and also T44A, F99S, Y145F, N146I, M153T or A, V163A, 1167T, S175G, S205T and N212K.

As used herein, the term "loop domain" refers to an amino acid sequence of an Aequorea-related fluorescent protein that connects the amino acids involved in the secondary structure of the eleven strands of the \(\pi\)-barrel or the central \(\pi\)-helix (residues 56-72) (see Fig. 1A and 1B).

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As used herein, the "fluorescent protein moiety" of a fluorescent protein is that portion of the amino acid sequence of a fluorescent protein which, when the amino acid sequence of the fluorescent protein substrate is optimally aligned with the amino acid sequence of a naturally occurring fluorescent protein, lies between the amino terminal and carboxy terminal amino acids, inclusive, of the amino acid sequence of the naturally occurring fluorescent protein.

It has been found that fluorescent proteins can be genetically fused to other target proteins and used as markers to identify the location and amount of the target protein produced. Accordingly, this invention provides fusion proteins comprising a fluorescent protein moiety and additional amino acid sequences. Such sequences can be, for example, up to about 15, up to about 50, up to about 150 or up to about 1000 amino acids long. The

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fusion proteins possess the ability to fluoresce when excited by electromagnetic radiation.

In one embodiment, the fusion protein comprises a polyhistidine tag to aid in purification of the protein.

# B. <u>Use Of The Crystal Structure Of Green Fluorescent Protein To Design</u> Mutants Having Altered Fluorescent Characteristics

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Using X-ray crystallography and computer processing, we have created a model of the crystal structure of Aequorea green fluorescent protein showing the relative location of the atoms in the molecule. This information is useful in identifying amino acids whose substitution alters fluorescent properties of the protein.

Fluorescent characteristics of Aequorea-related fluorescent proteins depend, in part, on the electronic environment of the chromophore. In general, amino acids that are within about 0.5 nm of the chromophore influence the electronic environment of the chromophore. Therefore, substitution of such amino acids can produce fluorescent proteins with altered fluorescent characteristics. In the excited state, electron density tends to shift from the phenolate towards the carbonyl end of the chromophore. Therefore, placement of increasing positive charge near the carbonyl end of the chromophore tends to decrease the energy of the excited state and cause a red-shift in the absorbance and emission wavelength maximum of the protein. Decreasing positive charge near the carbonyl end of the

Amino acids with charged (ionized D, E, K, and R), dipolar (H, N, Q, S, T, and uncharged D, E and K), and polarizable side groups (e.g., C, F, H, M, W and Y) are useful for altering the electronic environment of the chromophore, especially when they substitute an amino acid with an uncharged, nonpolar or non-polarizable side chain. In general, amino acids with polarizable side groups alter the electronic environment least, and, consequently, are expected to cause a comparatively smaller change in a fluorescent property. Amino acids with charged side groups alter the environment most, and, consequently, are expected to cause a comparatively larger change in a fluorescent property. However, amino acids with charged side groups are more likely to disrupt the structure of the protein and to prevent proper folding if buried next to the chromophore without any

additional solvation or salt bridging. Therefore charged amino acids are most likely to be tolerated and to give useful effects when they replace other charged or highly polar amino acids that are already solvated or involved in salt bridges. In certain cases, where substitution with a polarizable amino acid is chosen, the structure of the protein may make selection of a larger amino acid, e.g., W, less appropriate. Alternatively, positions occupied by amino acids with charged or polar side groups that are unfavorably oriented may be substituted with amino acids that have less charged or polar side groups. In another alternative, an amino acid whose side group has a dipole oriented in one direction in the protein can be substituted with an amino acid having a dipole oriented in a different direction.

More particularly, Table B lists several amino acids located within about 0.5 nm from the chromophore whose substitution can result in altered fluorescent characteristics. The table indicates, underlined, preferred amino acid substitutions at the indicated location to alter a fluorescent characteristic of the protein. In order to introduce such substitutions, the table also provides codons for primers used in site-directed mutagenesis involving amplification. These primers have been selected to encode economically the preferred amino acids, but they encode other amino acids as well, as indicated, or even a stop codon, denoted by Z. In introducing substitutions using such degenerate primers the most efficient strategy is to screen the collection to identify mutants with the desired properties and then sequence their DNA to find out which of the possible substitutions is responsible. Codons are shown in double-stranded form with sense strand above, antisense strand below. In nucleic acid sequences, R=(A or g); Y=(C or T); M=(A or C); K=(g or T); S=(g or C); W=(A or T); H=(A, T, or C); B=(g, T, or C); V=(g, A, or C);

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#### TABLE B

Original position and presumed role Change to Codon

L42 Aliphatic residue near C=N of chromophore CFHLQRWYZ 5YDS 3' 3'RHS 5'

V61 Aliphatic residue near central -CH= of chromophore FYHCLR YDC RHg

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	T62	. Almost directly above center of chromophore bridge $\underline{\text{AVFS}}$	KYC	MRg
5			DEHKNO	VAS BTS
			FYHCLR	YDC
10	V68	Aliphatic residue near carbonyl and G67	<u>FYH</u> L	RHg YWC RWg
	N121	Near C-N site of ring closure between T65 and G67 CFHLQ	RWYZ YDS	
15				RHS
•	Y145	Packs near tyrosine ring of chromophore	WCFL	TKS AMS
20			D <u>EHNKQ</u>	VAS BTS
	H148	H-bonds to phenolate oxygen	FYNI	wwc wwg
25			KQR	MRg KYC
	V150	Aliphatic residue near tyrosine ring of chromophore FYHL	YWC	RWg
30	F165	Packs near tyrosine ring	C <u>HO</u> R <u>WY</u> Z	YRS RYS
35	1167	Aliphatic residue near phenolate; 1167T has effects	FYHL	YWC RWg
	T203	H-bonds to phenolic oxygen of chromophore	<u>FH</u> LQR <u>WY</u> Z	YDS RHS
40	E222	Protonation regulates ionization of chromophore	нкио	MAS KTS

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Examples of amino acids with polar side groups that can be substituted with polarizable side groups include, for example, those in Table C.

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#### TABLE C

	Origina	al position and presumed role	Change to	Codon
5	Q69	Terminates chain of H-bonding waters	KREG	RRg YYC
10	Q94	H-bonds to carbonyl terminus of chromophore	<u>DEHKN</u> Q	VAS BTS
10	Q183	Bridges Arg96 and center of chromophore bridge	<u>HY</u>	YAC RTG
15			<u>EK</u>	RAg YTC
	N185	Part of H-bond network near carbonyl of chromophore	DEHNKQ	VAS BTS

In another embodiment, an amino acid that is close to a second amino acid within about 0.5 nm of the chromophore can, upon substitution, alter the electronic properties of the second amino acid, in turn altering the electronic environment of the chromphore. Table D presents two such amino acids. The amino acids, L220 and V224, are close to E222 and oriented in the same direction in the D pleated sheet.

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Original position and presumed role

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#### TABLE D

Change to

Codon

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30	L220	Packs next to Glu222; to make GFP pH sensitive	<u>Н</u> К <u>И</u> Р <u>Q</u> Т	MMS KKS
35	V224	Packs next to Glu222; to make GFP pH sensitive	<u>НКИРОТ</u>	MMS KKS
			C <u>FH</u> LQR <u>WY</u> Z	YDS

One embodiment of the invention includes a nucleic acid molecule comprising a nucleotide sequence encoding a functional engineered fluorescent protein whose amino acid sequence is substantially identical to the amino acid sequence of Aequorea green fluorescent protein (SEQ ID NO:2) and which differs from SEQ ID NO:2 by at least a substitution at Q69, wherein the functional engineered fluorescent protein has a different fluorescent property than Aequorea green fluorescent protein. Preferably, the substitution at Q69 is selected from the group of K, R, E and G. The Q69 substitution can be combined with other mutations to improve the properties of the protein, such as a functional mutation at S65.

One embodiment of the invention includes a nucleic acid molecule comprising a nucleotide sequence encoding a functional engineered fluorescent protein whose amino acid sequence is substantially identical to the amino acid sequence of Aequorea green fluorescent protein (SEQ ID NO:2) and which differs from SEQ ID NO:2 by at least a substitution at E222, but not including E222G, wherein the functional engineered fluorescent protein has a different fluorescent property than Aequorea green fluorescent protein. Preferably, the substitution at E222 is selected from the group of N and Q. The E222 substitution can be combined with other mutations to improve the properties of the protein, such as a functional mutation at F64.

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One embodiment of the invention includes a nucleic acid molecule comprising a nucleotide sequence encoding a functional engineered fluorescent protein whose amino acid sequence is substantially identical to the amino acid sequence of Aequorea green fluorescent protein (SEQ ID NO:2) and which differs from SEQ ID NO:2 by at least a substitution at Y145, wherein the functional engineered fluorescent protein has a different fluorescent property than Aequorea green fluorescent protein.

Preferably, the substitution at Y145 is selected from the group of W, C, F, L, E, H, K and Q.

The Y145 substitution can be combined with other mutations to improve the properties of the protein, such as a Y66.

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The invention also includes computer related embodiments, including computational methods of using the crystal coordinates for designing new fluorescent protein mutations and devices for storing the crystal data, including coordinates. For instance the invention includes a device comprising a storage device and, stored in the device, at least 10 atomic coordinates selected from the atomic coordinates listed in Figs. 5-1 to 5-28. More coordinates can be storage depending of the complexity of the calculations or the objective of using the coordinates (e.g. about 100, 1,000, or more coordinates). For example, larger numbers of coordinates will be desirable for more detailed representations of fluorescent protein structure. Typically, the storage device is a computer readable device that stores code that it receives as input the atomic coordinates. Although, other storage meand as known in the art are contemplated. The computer readable device can be a floppy disk or a hard drive.

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#### C. Production Of Long Wavelength Engineered Fluorescent Proteins

Recombinant production of a fluorescent protein involves expressing a nucleic acid molecule having sequences that encode the protein.

In one embodiment, the nucleic acid encodes a fusion protein in which a single polypeptide includes the fluorescent protein moiety within a longer polypeptide. The longer polypeptide can include a second functional protein, such as FRET partner or a protein having a second function (e.g., an enzyme, antibody or other binding protein). Nucleic acids that encode fluorescent proteins are useful as starting materials.

The fluorescent proteins can be produced as fusion proteins by recombinant DNA technology. Recombinant production of fluorescent proteins involves expressing nucleic acids having sequences that encode the proteins. Nucleic acids encoding fluorescent proteins can be obtained by methods known in the art. Fluorescent proteins can be made by site-specific mutagenesis of other nucleic acids encoding fluorescent proteins, or by random mutagenesis caused by increasing the error rate of PCR of the original polynucleotide with 0.1 mM MnCl, and unbalanced nucleotide concentrations. See, e.g., U.S. patent application 08/337,915, filed November 10, 1994 or International application PCT/US95/14692, filed 11/10/95. The nucleic acid encoding a green fluorescent protein can be isolated by polymerase chain reaction of cDNA from A. victoria using primers based on the DNA sequence of A. victoria green fluorescent protein, as presented in Fig. 3. PCR methods are described in, for example, U.S. Pat. No. 4,683,195; Mullis et al. (1987) Cold Spring Harbor Symp. Quant. Biol. 51:263; and Erlich, ed., PCR Technology, (Stockton Press, NY, 1989).

The construction of expression vectors and the expression of genes in transfected cells involves the use of molecular cloning techniques also well known in the art. Sambrook et al., Molecular Cloning — A Laboratory Manual, Cold Spring Harbor Laboratory, Cold Spring Harbor, NY, (1989) and Current Protocols in Molecular Biology, F.M. Ausubel et al., eds., (Current Protocols, a joint venture between Greene Publishing Associates, Inc. and John Wiley & Sons, Inc.). The expression vector can be adapted for function in prokaryotes or eukaryotes by inclusion of appropriate promoters, replication sequences, markers, etc.

Nucleic acids used to transfect cells with sequences coding for expression of the polypeptide of interest generally will be in the form of an expression vector including

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expression control sequences operatively linked to a nucleotide sequence coding for expression of the polypeptide. As used, the term "nucleotide sequence coding for expression of" a polypeptide refers to a sequence that, upon transcription and translation of mRNA, produces the polypeptide. This can include sequences containing, e.g., introns. Expression control sequences are operatively linked to a nucleic acid sequence when the expression control sequences control and regulate the transcription and, as appropriate, translation of the nucleic acid sequence. Thus, expression control sequences can include appropriate promoters, enhancers, transcription terminators, a start codon (i.e., ATG) in front of a protein-encoding gene, splicing signals for introns, maintenance of the correct reading frame of that sene to permit proper translation of the mRNA, and stop codons.

Methods which are well known to those skilled in the art can be used to construct expression vectors containing the fluorescent protein coding sequence and appropriate transcriptional/translational control signals. These methods include in vitro recombinant DNA techniques, synthetic techniques and in vivo recombination/genetic recombination. (See, for example, the techniques described in Maniatis, et al., Molecular Cloning A Laboratory Manual, Cold Spring Harbor Laboratory, N.Y., 1989).

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Transformation of a host cell with recombinant DNA may be carried out by conventional techniques as are well known to those skilled in the art. Where the host is prokaryotic, such as E. coli, competent cells which are capable of DNA uptake can be prepared from cells harvested after exponential growth phase and subsequently treated by the CaCl<sub>2</sub> method by procedures well known in the art. Alternatively, MgCl<sub>2</sub> or RbCl can be used. Transformation can also be performed after forming a protoplast of the host cell or by electroporation.

When the host is a eukaryote, such methods of transfection of DNA as calcium phosphate co-precipitates, conventional mechanical procedures such as microinjection, electroporation, insertion of a plasmid encased in liposomes, or virus vectors may be used. Eukaryotic cells can also be cotransfected with DNA sequences encoding the fusion polypeptide of the invention, and a second foreign DNA molecule encoding a selectable phenotype, such as the herpes simplex thymidine kinase gene. Another method is to use a eukaryotic viral vector, such as simian virus 40 (SV40) or bovine papilloma virus, to transiently infect or transform eukaryotic cells and express the protein. (Eukaryotic Viral

Vectors, Cold Spring Harbor Laboratory, Gluzman ed., 1982). Preferably, a eukaryotic host is utilized as the host cell as described herein.

Techniques for the isolation and purification of either microbially or eukaryotically expressed polypeptides of the invention may be by any conventional means such as, for example, preparative chromatographic separations and immunological separations such as those involving the use of monoclonal or polyclonal antibodies or antigen.

In one embodiment recombinant fluorescent proteins can be produced by expression of nucleic acid encoding for the protein in *E. coli. Aequorea*-related fluorescent proteins are best expressed by cells cultured between about 15 \(\times\) C and 30 \(\times\) C but higher temperatures (e.g. 37 \(\times\) C) are possible. After synthesis, these enzymes are stable at higher temperatures (e.g., 37 \(\times\) C) and can be used in assays at those temperatures.

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A variety of host-expression vector systems may be utilized to express fluorescent protein coding sequence. These include but are not limited to microorganisms such as bacteria transformed with recombinant bacteriophage DNA, plasmid DNA or cosmid DNA expression vectors containing a fluorescent protein coding sequence; yeast transformed with recombinant yeast expression vectors containing the fluorescent protein coding sequence; plant cell systems infected with recombinant virus expression vectors (e.g., cauliflower mosaic virus, CaMV; tobacco mosaic virus, TMV) or transformed with recombinant plasmid expression vectors (e.g., Ti plasmid) containing a fluorescent protein coding sequence; insect cell systems infected with recombinant virus expression vectors (e.g., baculovirus) containing a fluorescent protein coding sequence; or animal cell systems infected with recombinant virus expression vectors (e.g., retroviruses, adenovirus, vaccinia virus) containing a fluorescent protein coding sequence; or animal cell systems engineered for stable expression.

Depending on the host/vector system utilized, any of a number of suitable transcription and translation elements, including constitutive and inducible promoters, transcription enhancer elements, transcription terminators, etc. may be used in the expression vector (see, e.g., Bitter, et al., Methods in Enzymology 153:516-544, 1987). For example, when cloning in bacterial systems, inducible promoters such as pL of bacteriophage D, plac, ptrp, ptac (ptrp-lac hybrid promoter) and the like may be used. When cloning in mammalian cell systems, promoters derived from the genome of mammalian cells (e.g., metallothionein promoter) or from mammalian viruses (e.g., the

retrovirus long terminal repeat; the adenovirus late promoter; the vaccinia virus 7.5K promoter) may be used. Promoters produced by recombinant DNA or synthetic techniques may also be used to provide for transcription of the inserted fluorescent protein coding sequence.

In bacterial systems a number of expression vectors may be advantageously selected depending upon the use intended for the fluorescent protein expressed. For example, when large quantities of the fluorescent protein are to be produced, vectors which direct the expression of high levels of fusion protein products that are readily purified may be desirable. Those which are engineered to contain a cleavage site to aid in recovering fluorescent protein are preferred.

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In yeast, a number of vectors containing constitutive or inducible promoters may be used. For a review see, Current Protocols in Molecular Biology, Vol. 2, Ed. Ausubel, et al., Greene Publish. Assoc. & Wiley Interscience, Ch. 13, 1988; Grant, et al., Expression and Secretion Vectors for Yeast, in Methods in Enzymology, Eds. Wu & Grossman, 31987, Acad. Press, N.Y., Vol. 153, pp.516-544, 1987; Glover, DNA Cloning, Vol. II, IRL Press, Wash., D.C., Ch. 3, 1986; and Bitter, Heterologous Gene Expression in Yeast, Methods in Enzymology, Eds. Berger & Kimmel, Acad. Press, N.Y., Vol. 152, pp. 673-684, 1987; and The Molecular Biology of the Yeast Saccharomyces, Eds. Strathern et al., Cold Spring Harbor Press, Vols. I and II, 1982. A constitutive yeast promoter such as ADH or LEU2 or an inducible promoter such as GAL may be used (Cloning in Yeast, Ch. 3, R. Rothstein In: DNA Cloning Vol.11, A Practical Approach, Ed. DM Glover, RRL Press, Wash., D.C., 1986). Alternatively, vectors may be used which promote integration of foreign DNA sequences into the yeast chromosome.

In cases where plant expression vectors are used, the expression of a fluorescent protein coding sequence may be driven by any of a number of promoters. For example, viral promoters such as the 35S RNA and 19S RNA promoters of CaMV (Brisson, et al., Nature 310:511-514, 1984), or the coat protein promoter to TMV (Takamatsu, et al., EMBO J. 6:307-311, 1987) may be used; alternatively, plant promoters such as the small subunit of RUBISCO (Coruzzi, et al., 1984, EMBO J. 3:1671-1680; Broglie, et al., Science 224:838-843, 1984); or heat shock promoters, e.g., soybean hsp17.5-E or hsp17.3-B (Gurley, et al., Mol. Cell. Biol. 6:559-565, 1986) may be used. These constructs can be introduced into plant cells using Ti plasmids, Ri plasmids, plant virus vectors, direct DNA transformation,

microinjection, electroporation, etc. For reviews of such techniques see, for example, Weissbach & Weissbach, Methods for Plant Molecular Biology, Academic Press, NY, Section VIII, pp. 421-463, 1988; and Grierson & Corey, Plant Molecular Biology, 2d Ed., Blackie. London. Ch. 7-9, 1988.

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An alternative expression system which could be used to express fluorescent protein is an insect system. In one such system, Autographa californica nuclear polyhedrosis virus (AcNPV) is used as a vector to express foreign genes. The virus grows in Spodoptera frugiperda cells. The fluorescent protein coding sequence may be cloned into non-essential regions (for example, the polyhedrin gene) of the virus and placed under control of an AcNPV promoter (for example the polyhedrin promoter). Successful insertion of the fluorescent protein coding sequence will result in inactivation of the polyhedrin gene and production of non-occluded recombinant virus (i.e., virus lacking the proteinaceous coat coded for by the polyhedrin gene). These recombinant viruses are then used to infect Spodoptera frugiperda cells in which the inserted gene is expressed, see Smith, et al., J. Viol. 46:584, 1983; Smith, U.S. Patent No. 4,215,051.

Eukaryotic systems, and preferably mammalian expression systems, allow for proper post-translational modifications of expressed mammalian proteins to occur. Eukaryotic cells which possess the cellular machinery for proper processing of the primary transcript, glycosylation, phosphorylation, and, advantageously secretion of the gene product should be used as host cells for the expression of fluorescent protein. Such host cell lines may include but are not limited to CHO, VERO, BHK, HeLa, COS, MDCK, Jurkat. HEK-293, and WI38.

Mammalian cell systems which utilize recombinant viruses or viral elements to direct expression may be engineered. For example, when using adenovirus expression vectors, the fluorescent protein coding sequence may be ligated to an adenovirus transcription/translation control complex, e.g., the late promoter and tripartite leader sequence. This chimeric gene may then be inserted in the adenovirus genome by in vitro or in vivo recombination. Insertion in a non-essential region of the viral genome (e.g., region E1 or E3) will result in a recombinant virus that is viable and capable of expressing the fluorescent protein in infected hosts (e.g., see Logan & Shenk, Proc. Natl. Acad. Sci. USA, 81: 3655-3659, 1984). Alternatively, the vaccinia virus 7.5K promoter may be used. (e.g., see, Mackett, et al., Proc. Natl. Acad. Sci. USA, 79: 7415-7419, 1982: Mackett, et al., J.

Virol. 49: 857-864, 1984; Panicali, et al., Proc. Natl. Acad. Sci. USA 79: 4927-4931, 1982). Of particular interest are vectors based on bovine papilloma virus which have the ability to replicate as extrachromosomal elements (Sarver, et al., Mol. Cell. Biol. 1: 486, 1981). Shortly after entry of this DNA into mouse cells, the plasmid replicates to about 100 to 200 copies per cell. Transcription of the inserted cDNA does not require integration of the plasmid into the host's chromosome, thereby yielding a high level of expression. These vectors can be used for stable expression by including a selectable marker in the plasmid, such as the neo gene. Alternatively, the retroviral genome can be modified for use as a vector capable of introducing and directing the expression of the fluorescent protein gene in host cells (Cone & Mulligan, Proc. Natl. Acad. Sci. USA, 81:6349-6353, 1984). High level expression may also be achieved using inducible promoters, including, but not limited to, the metallothionine IIA promoter and heat shock promoters.

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The invention can also include a localization sequence, such as a nuclear localization sequence, an endoplasmic reticulum localization sequence, a peroxisome localization sequence, a mitochondrial localization sequence, or a localized protein.

Localization sequences can be targeting sequences which are described, for example, in "Protein Targeting", chapter 35 of Stryer, L., Biochemistry (4th ed.). W.H. Freeman, 1995. The localization sequence can also be a localized protein. Some important localization sequences include those targeting the nucleus (KKKRK), mitochondrion (amino terminal MLRTSSLFTRRVQPSLFRNILRLQST-), endoplasmic reticulum (KDEL at C-terminus, assuming a signal sequence present at N-terminus), peroxisome (SKF at C-terminus), prenylation or insertion into plasma membrane (CaaX, CC, CXC, or CCXX at C-terminus), cytoplasmic side of plasma membrane (fusion to SNAP-25), or the Golgi apparatus (fusion to furin).

For long-term, high-yield production of recombinant proteins, stable expression is preferred. Rather than using expression vectors which contain viral origins of replication, host cells can be transformed with the fluorescent protein cDNA controlled by appropriate expression control elements (e.g., promoter, enhancer, sequences, transcription terminators, polyadenylation sites, etc.), and a selectable marker. The selectable marker in the recombinant plasmid confers resistance to the selection and allows cells to stably integrate the plasmid into their chromosomes and grow to form foci which in turn can be cloned and expanded into cell lines. For example, following the introduction of foreign DNA,

engineered cells may be allowed to grow for 1-2 days in an enriched media, and then are switched to a selective media. A number of selection systems may be used, including but not limited to the herpes simplex virus thymidine kinase (Wigler, et al., Cell, 11: 223, 1977), hypoxanthine-guanine phosphoribosyltransferase (Szybalska & Szybalski, Proc. Natl. Acad. Sci. USA, 48:2026, 1962), and adenine phosphoribosyltransferase (Lowy, et al., Cell, 22: 817, 1980) genes can be employed in tk', hgprt or aprt cells respectively. Also, antimetabolite resistance can be used as the basis of selection for dhfr. which confers resistance to methotrexate (Wigler, et al., Proc. Natl. Acad. Sci. USA, 77: 3567, 1980; O'Hare, et al., Proc. Natl. Acad. Sci. USA, 8: 1527, 1981); gpt, which confers resistance to mycophenolic acid (Mulligan & Berg, Proc. Natl. Acad. Sci. USA, 78: 2072, 1981; neo, which confers resistance to the aminoglycoside G-418 (Colberre-Garapin, et al., J. Mol. Biol., 150:1, 1981); and hygro, which confers resistance to hygromycin (Santerre, et al., Gene, 30: 147, 1984) genes. Recently, additional selectable genes have been described, namely trpB, which allows cells to utilize indole in place of tryptophan; hisD, which allows cells to utilize histinol in place of histidine (Hartman & Mulligan, Proc. Natl. Acad. Sci. USA, 85:8047, 1988); and ODC (ornithine decarboxylase) which confers resistance to the ornithine decarboxylase inhibitor, 2-(difluoromethyl)-DL-ornithine, DFMO (McConlogue L., In: Current Communications in Molecular Biology, Cold Spring Harbor Laboratory, ed., 1987).

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DNA sequences encoding the fluorescence protein polypeptide of the invention can be expressed in vitro by DNA transfer into a suitable host cell. "Host cells" are cells in which a vector can be propagated and its DNA expressed. The term also includes any progeny of the subject host cell. It is understood that all progeny may not be identical to the parental cell since there may be mutations that occur during replication. However, such progeny are included when the term "host cell" is used. Methods of stable transfer, in other words when the foreign DNA is continuously maintained in the host, are known in the art.

The expression vector can be transfected into a host cell for expression of the recombinant nucleic acid. Host cells can be selected for high levels of expression in order to purify the fluorescent protein fusion protein. *E. coli* is useful for this purpose.

Alternatively, the host cell can be a prokaryotic or eukaryotic cell selected to study the activity of an enzyme produced by the cell. In this case, the linker pentide is selected to

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include an amino acid sequence recognized by the protease. The cell can be, e.g., a cultured cell or a cell in vivo.

A primary advantage of fluorescent protein fusion proteins is that they are prepared by normal protein biosynthesis, thus completely avoiding organic synthesis and the requirement for customized unnatural amino acid analogs. The constructs can be expressed in E. coli in large scale for in vitro assays. Purification from bacteria is simplified when the sequences include polyhistidine tags for one-step purification by nickel-chelate chromatography. Alternatively, the substrates can be expressed directly in a desired host cell for assays in situ.

In another embodiment, the invention provides a transgenic non-human animal that expresses a nucleic acid sequence which encodes the fluorescent protein.

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The "non-human animals" of the invention comprise any non-human animal having nucleic acid sequence which encodes a fluorescent protein. Such non-human animals include vertebrates such as rodents, non-human primates, sheep, dog, cow, pig, amphibians, and reptiles. Preferred non-human animals are selected from the rodent family including rat and mouse, most preferably mouse. The "transgenic non-human animals" of the invention are produced by introducing "transgenes" into the germline of the non-human animal. Embryonal target cells at various developmental stages can be used to introduce transgenes. Different methods are used depending on the stage of development of the embryonal target cell. The zygote is the best target for micro-injection. In the mouse, the male pronucleus reaches the size of approximately 20 micrometers in diameter which allows reproducible injection of 1-2 pl of DNA solution. The use of zygotes as a target for gene transfer has a major advantage in that in most cases the injected DNA will be incorporated into the host gene before the first cleavage (Brinster et al., Proc. Natl. Acad. Sci. USA 82:4438-4442, 1985). As a consequence, all cells of the transgenic non-human animal will carry the incorporated transgene. This will in general also be reflected in the efficient transmission of the transgene to offspring of the founder since 50% of the germ cells will harbor the transgene. Microinjection of zygotes is the preferred method for incorporating transgenes in practicing the invention.

The term "transgenic" is used to describe an animal which includes exogenous genetic material within all of its cells. A "transgenie" animal can be produced by cross-breeding two chimeric animals which include exogenous genetic material within cells used

in reproduction. Twenty-five percent of the resulting offspring will be transgenic *i.e.*, animals which include the exogenous genetic material within all of their cells in both alleles. 50% of the resulting animals will include the exogenous genetic material within one allele and 25% will include no exogenous genetic material.

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Retroviral infection can also be used to introduce transgene into a non-human animal. The developing non-human embryo can be cultured in vitro to the blastocyst stage. During this time, the blastomeres can be targets for retro viral infection (Jaenich, R., Proc. Natl. Acad. Sci USA 73:1260-1264, 1976). Efficient infection of the blastomeres is obtained by enzymatic treatment to remove the zona pellucida (Hogan, et al. (1986) in Manipulating the Mouse Embryo, Cold Spring Harbor Laboratory Press, Cold Spring Harbor, N.Y.). The viral vector system used to introduce the transgene is typically a replication-defective retro virus carrying the transgene (Jahner, et al., Proc. Natl. Acad. Sci. USA 82:6927-6931, 1985; Van der Putten, et al., Proc. Natl. Acad. Sci USA 82:6148-6152, 1985). Transfection is easily and efficiently obtained by culturing the blastomeres on a monolayer of virus-producing cells (Van der Putten, supra; Stewart, et al., EMBO J. 6:383-388, 1987). Alternatively, infection can be performed at a later stage. Virus or virus-producing cells can be injected into the blastocoele (D. Jahner et al., Nature 298:623-628, 1982). Most of the founders will be mosaic for the transgene since incorporation occurs only in a subset of the cells which formed the transgenic nonhuman animal. Further, the founder may contain various retro viral insertions of the transgene at different positions in the genome which generally will segregate in the offspring. In addition, it is also possible to introduce transgenes into the germ line, albeit with low efficiency, by intrauterine retro viral infection of the midgestation embryo (D. Jahner et al., supra).

A third type of target cell for transgene introduction is the embryonal stem cell (ES). ES cells are obtained from pre-implantation embryos cultured in vitro and fused with embryos (M. J. Evans et al. Nature 292:154-156, 1981; M.O. Bradley et al., Nature 309: 255-258, 1984; Gossler, et al., Proc. Natl. Acad. Sci. USA 83: 9065-9069, 1986; and Robertson et al., Nature 322:445-448, 1986). Transgenes can be efficiently introduced into the ES cells by DNA transfection or by retro virus-mediated transduction. Such transformed ES cells can thereafter be combined with blastocysts from a nonhuman animal. The ES cells thereafter colonize the embryo and contribute to the germ line of the resulting chimeric animal. (For review see Jaenisch, R., Science 240: 1468-1474, 1988).

"Transformed" means a cell into which (or into an ancestor of which) has been introduced, by means of recombinant nucleic acid techniques, a heterologous nucleic acid molecule. "Heterologous" refers to a nucleic acid sequence that either originates from another species or is modified from either its original form or the form primarily expressed in the cell.

"Transgene" means any piece of DNA which is inserted by artifice into a cell, and becomes part of the genome of the organism (i.e., either stably integrated or as a stable extrachromosomal element) which develops from that cell. Such a transgene may include a gene which is partly or entirely heterologous (i.e., foreign) to the transgenic organism, or may represent a gene homologous to an endogenous gene of the organism. Included within this definition is a transgene created by the providing of an RNA sequence which is transcribed into DNA and then incorporated into the genome. The transgenes of the invention include DNA sequences which encode which encodes the fluorescent protein which may be expressed in a transgenic non-human animal. The term "transgenic" as used herein additionally includes any organism whose genome has been altered by in vitro manipulation of the early embryo or fertilized egg or by any transgenic technology to induce a specific gene knockout. The term "gene knockout" as used herein, refers to the targeted disruption of a gene in vivo with complete loss of function that has been achieved by any transgenic technology familiar to those in the art. In one embodiment, transgenic animals having gene knockouts are those in which the target gene has been rendered nonfunctional by an insertion targeted to the gene to be rendered non-functional by homologous recombination. As used herein, the term "transgenic" includes any transgenic technology familiar to those in the art which can produce an organism carrying an introduced transgene or one in which an endogenous gene has been rendered non-functional or "knocked out."

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#### III USES OF ENGINEERED FLUORESCENT PROTEINS

The proteins of this invention are useful in any methods that employ fluorescent proteins.

The engineered fluorescent proteins of this invention are useful as fluorescent markers in the many ways fluorescent markers already are used. This includes, for example, coupling engineered fluorescent proteins to antibodies, nucleic acids or other recentors for use in detection assays, such as immunoassays or hybridization assays.

The engineered fluorescent proteins of this invention are useful to track the movement of proteins in cells. In this embodiment, a nucleic acid molecule encoding the fluorescent protein is fused to a nucleic acid molecule encoding the protein of interest in an expression vector. Upon expression inside the cell, the protein of interest can be localized based on fluorescence. In another version, two proteins of interest are fused with two engineered fluorescent proteins having different fluorescent characteristics.

The engineered fluorescent proteins of this invention are useful in systems to detect induction of transcription. In certain embodiments, a nucleotide sequence encoding the engineered fluorescent protein is fused to expression control sequences of interest and the expression vector is transfected into a cell. Induction of the promoter can be measured by detecting the expression and/or quantity of fluorescence. Such constructs can be used used to follow signaling pathways from receptor to promoter.

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The engineered fluorescent proteins of this invention are useful in applications involving FRET. Such applications can detect events as a function of the movement of fluorescent donors and acceptor towards or away from each other. One or both of the donor/acceptor pair can be a fluorescent protein. A preferred donor and receptor pair for FRET based assays is a donor with a T203I mutation and an acceptor with the mutation T203X, wherein X is an aromatic amino acid-39, especially T203Y, T203W, or T203H. In a particularly useful pair the donor contains the following mutations: S72A, K79R, Y145F, M153A and T203I (with a excitation peak of 395 nm and an emission peak of 511 nm) and the acceptor contains the following mutations S65G, S72A, K79R, and \*T203Y. This particular pair provides a wide separation between the excitation and emission peaks of the donor and provides good overlap between the donor emission spectrum and the acceptor excitation spectrum. Other red-shifted mutants, such as those described herein, can also be used as the acceptor in such a pair.

In one aspect, FRET is used to detect the cleavage of a substrate having the donor and acceptor coupled to the substrate on opposite sides of the cleavage site. Upon cleavage of the substrate, the donor/acceptor pair physically separate, eliminating FRET. Assays involve contacting the substrate with a sample, and determining a qualitative or quantitative change in FRET. In one embodiment, the engineered fluorescent protein is used in a substrate for  $\Box$ -lactamase. Examples of such substrates are described in United States patent applications 08/407.544, filed March 20, 1995 and International Application

PCT/US96/04059, filed March 20, 1996. In another embodiment, an engineered fluorescent protein donor/acceptor pair are part of a fusion protein coupled by a peptide having a proteolytic cleavage site. Such tandem fluorescent proteins are described in United States patent application 08/594,575, filed January 31, 1996.

In another aspect, FRET is used to detect changes in potential across a membrane. A donor and acceptor are placed on opposite sides of a membrane such that one translates across the membrane in response to a voltage change. This creates a measurable FRET. Such a method is described in United States patent application 08/481,977, filed June 7, 1995 and International Application PCT/US96/09652, filed June 6, 1996.

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The engineered protein of this invention are useful in the creation of fluorescent substrates for protein kinases. Such substrates incorporate an amino acid sequence recognizable by protein kinases. Upon phosphorylation, the engineered fluorescent protein undergoes a change in a fluorescent property. Such substrates are useful in detecting and measuring protein kinase activity in a sample of a cell, upon transfection and expression of the substrate. Preferably, the kinase recognition site is placed within about 20 amino acids of a terminus of the engineered fluorescent protein. The kinase recognition site also can be placed in a loop domain of the protein. (See, e.g. Figure 1B.) Methods for making fluorescent substrates for protein kinases are described in United States patent application 08/680,877, filed July 16, 1996.

A protease recognition site also can be introduced into a loop domain. Upon cleavage, fluorescent property changes in a measurable fashion.

The invention also includes a method of identifying a test chemical. Typically, the method includes contacting a test chemical a sample containing a biological entity labeled with a functional, engineered fluorescent protein or a polynucleotide encoding said functional, engineered fluorescent protein. By monitoring fluorescence (i.e. a fluorescent property) from the sample containing the functional engineered fluorescent protein it can be determined whether a test chemical is active. Controls can be included to insure the specificity of the signal. Such controls include measurements of a fluorescent property in the absence of the test chemical, in the presence of a chemical with an expected activity (e.g., a known modulator) or engineered controls (e.g., absence of engineered fluorescent protein, absence of engineered fluorescent protein polynucleotide or the absence of operably linkage of the engineered fluorescent protein).

The fluorescence in the presence of a test chemical can be greater or less than in the absence of said test chemical. For instance if the engineered fluorescent protein is used a reporter of gene expression, the test chemical may up or down regulate gene expression. For such types of screening, the polynucleotide encoding the functional, engineered fluorescent protein is operatively linked to a genomic polynucleotide or a re. Alternatively, the functional, engineered fluorescent protein is fused to second functional protein. This embodiment can be used to track localization of the second protein or to track protein-protein interactions using energy transfer.

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### IV. PROCEDURES

Fluorescence in a sample is measured using a fluorimeter. In general, excitation radiation from an excitation source having a first wavelength, passes through excitation optics. The excitation optics cause the excitation radiation to excite the sample.

In response, fluorescent proteins in the sample emit radiation which has a wavelength that is different from the excitation wavelength. Collection optics then collect the emission from the sample. The device can include a temperature controller to maintain the sample at a specific temperature while it is being scanned. According to one embodiment, a multi-axis translation stage moves a microtiter plate holding a plurality of samples in order to position different wells to be exposed. The multi-axis translation stage, temperature controller, autofocusing feature, and electronics associated with imaging and data collection can be managed by an appropriately programmed digital computer. The computer also can

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transform the data collected during the assay into another format for presentation. This process can be miniaturized and automated to enable screening many thousands of compounds.

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Methods of performing assays on fluorescent materials are well known in the art and are described in, e.g., Lakowicz, J.R., Principles of Fluorescence Spectroscopy, New York: Plenum Press (1983); Herman, B., Resonance energy transfer microscopy, in: Fluorescence Microscopy of Living Cells in Culture, Part B, Methods in Cell Biology, vol. 30, ed. Taylor, D.L. & Wang, Y.-L., San Diego: Academic Press (1989), pp. 219-243; Turro, N.J., Modern Molecular Photochemistry, Menlo Park: Benjamin/Cummings Publishing Col, Inc. (1978), pp. 296-361.

The following examples are provided by way of illustration, not by way of limitation.

#### EXAMPLES

As a step in understanding the properties of GFP, and to aid in the tailoring of GFPs with altered characteristics, we have determined the three dimensional structure at 1.9 Å resolution of the S65T mutant (R. Heim et al. Nature 373:664-665 (1995)) of A. victoria GFP. This mutant also contains the ubiquitous Q80R substitution, which accidentally occurred in the early distribution of the GFP cDNA and is not known to have any effect on the protein properties (M. Chalfie et al. Science 263:802-805 (1994)).

Histidine-tagged S65T GFP (R. Heim et al. *Nature* 373:664-665 (1995)) was overexpressed in JM109/pRSET<sub>a</sub> in 41 YT broth plus ampicillin at 37□, 450 rpm and 5 l/min air flow. The temperature was reduced to 25□ at A<sub>995</sub> = 0.3, followed by induction with 1mM isopropylthiogalactoside for 5h. Cell paste was stored at -80□ overnight, then was resuspended in 50 mM HEPES pH 7.9, 0.3 M NaCl, 5 mM 2-mercaptoethanol, 0.1 mM phenylmethyl-sulfonylfluoride (PMSF), passed once through a French press at 10,000 psi, then centrifuged at 20 K rpm for 45 min. The supernatant was applied to a Ni-NTA-agarose column (Qiagen), followed by a wash with 20 mM imidazole, then eluted with 100 mM imidazole. Green fractions were pooled and subjected to chymotryptic (Sigma) proteolysis (1:50 w/w) for 22 h at RT. After addition of 0.5 mM PMSF, the digest was reapplied to the

Ni column. N-terminal sequencing verified the presence of the correct N-terminal methionine. After dialysis against 20 mM HEPES, pH 7.5 and concentration to  $A_{400}$  = 20, rod-shaped crystals were obtained at RT in hanging drops containing 5  $\square$ 1 protein and 5  $\square$ 1 well solution, 22-26% PEG 4000 (Serva), 50 mM HEPES pH 8.0-8.5, 50 mM MgCl<sub>2</sub> and 10 mM 2-mercapto-ethanol within 5 days. Crystals were 0.05 mm across and up to 1.0 mm long. The space group is P2,2,2, with a = 51.8, b = 62.8, c = 70.7 Å, Z=4. Two crystal forms of wild-type GFP, unrelated to the present form, have been described by M. A. Perrozo, K. B. Ward, R. B. Thompson, & W. W. Ward. *J. Biol. Chem.* 203, 7713-7716 (1988).

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The structure of GFP was determined by multiple isomorphous replacement and anomalous scattering (Table E), solvent flattening, phase combination and crystallographic refinement. The most remarkable feature of the fold of GFP is an eleven stranded \( \beta\)-barrel wrapped around a single central helix (Fig. 1A and 1B), where each strand consists of approximately 9-13 residues. The barrel forms a nearly perfect cylinder 42 Å long and 24 Å in diameter. The N-terminal half of the polypeptide comprises three antiparallel strands, the central helix, and then 3 more anti-parallel strands, the latter of which (residues 118-123) is parallel to the N-terminal strand (residues 11-23). The polypeptide backbone then crosses the "bottom" of the molecule to form the second half of the barrel in a five-strand Greek Key motif. The top end of the cylinder is capped by three short, distorted helical segments, while one short, very distorted helical segment caps the bottom of the cylinder. The main-chain hydrogen bonding lacing the surface of the cylinder very likely accounts for the unusual stability of the protein towards denaturation and proteolysis. There are no large segments of the polypeptide that could be excised while preserving the intactness of the shell around the chromophore. Thus it would seem difficult to re-engineer GFP to reduce its molecular weight (J. Dopf & T.M. Horiagon Gene 173:39-43 (1996)) by a large percentage.

The p-hydroxybenzylideneimidazolidinone chromophore (C. W. Cody et al. Biochemistry 32:1212-1218 (1993)) is completely protected from bulk solvent and centrally located in the molecule. The total and presumably rigid encapsulation is probably responsible for the small Stokes' shift (i.e. wavelength difference between excitation and emission maxima), high quantum yield of fluorescence, inability of O<sub>2</sub> to quench the excited state (B.D. Nageswara Rao et al. Biophys. J. 32:630-632 (1980)), and resistance of the chromophore to titration of the external pH (W. W. Ward. Bioluminescence and Chemiluminescence (M. A. DeLuca and W. D. McElroy, eds) Academic Press pp. 235-242 (1981); W. W. Ward & S. H. Bokman, Biochemistry 21:4535-4540 (1982); W. W. Ward et al. Photochem. Photobiol. 35:803-808 (1982)). It also allows one to rationalize why fluorophore formation should be a spontaneous intramolecular process (R. Heim et al. Proc. Natl. Acad. Sci. USA 91:12501-12504 (1994)), as it is difficult to imagine how an enzyme could gain access to the substrate. The plane of the chromophore is roughly perpendicular (60□) to the symmetry axis of the surrounding barrel. One side of the chromophore faces a surprisingly large cavity, that occupies a volume of approximately 135 Å3 (B. Lee & F. M. Richards, J. Mol. Biol. 55:379-400 (1971)). The atomic radii were those of Lee & Richards, calculated using the program MS with a probe radius of 1.4 Å. (M. L. Connolly, Science 221:709-713 (1983)). The cavity does not open out to bulk solvent. Four water molecules are located in the cavity, forming a chain of hydrogen bonds linking the buried side chains of Glu222 and Gln69. Unless occupied, such a large cavity would be expected to de-stabilize the protein by several kcal/mol (S. J. Hubbard et al., Protein Engineering 7:613-626 (1994); A. E. Eriksson et al. Science 255:178-183 (1992)). Part of the volume of the cavity might be the consequence of the compaction resulting from cyclization and dehydration reactions. The cavity might also temporarily accommodate the oxidant, most likely O<sub>2</sub> (A. B. Cubitt et al. Trends Biochem. Sci. 20:448-455 (1995); R. Heim et al. Proc. Natl. Acad. Sci. USA 91:12501-12504 (1994); S. Inouye & F.I. Tsuji. FEBS Lett. 351:211-214 (1994)), that dehydrogenates the □-□ bond of Tyr66. The chromophore, cavity, and side chains that contact the chromophore are shown in Figure 2A and a portion of the final electron density

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map in this vicinity in 2B.

The opposite side of the chromophore is packed against several aromatic and polar side chains. Of particular interest is the intricate network of polar interactions with the chromophore (Fig. 2C). His <sup>148</sup>, Thr<sup>260</sup> and Ser<sup>260</sup> form hydrogen bonds with the phenolic hydroxyl; Arg<sup>86</sup> and Gin<sup>84</sup> interact with the carbonyl of the imidazolidinone ring and Ghu<sup>222</sup> forms a hydrogen bond with the side chain of Thr<sup>85</sup>. Additional polar interactions, such as hydrogen bonds to Arg<sup>86</sup> from the carbonyl of Thr<sup>82</sup>, and the side-chain carbonyl of Gln<sup>143</sup>, presumably stabilize the buried Arg<sup>86</sup> in its protonated form. In turn, this buried charge suggests that a partial negative charge resides on the carbonyl oxygen of the imidazolidinone rine of the deprotonated fluorophore, as has previously been suggested (W.

W. Ward. Bioluminescence and Chemiluminescence (M. A. DeLuca and W. D. McElroy, eds) Academic Press pp. 235-242 (1981); W. W. Ward & S. H. Bokman. Biochemistry 21:4535-4540 (1982); W. W. Ward et al. Photochem. Photobiol. 35:803-808 (1982)). Arg<sup>36</sup> is likely to be essential for the formation of the fluorophore, and may help catalyze the initial ring closure. Finally, Tyr<sup>143</sup> shows a typical stabilizing edge-face interaction with the benzyl ring. Trp<sup>57</sup>, the only tryptophan in GFP, is located 13 Å to 15 Å from the chromophore and the long axes of the two ring systems are nearly parallel. This indicates that efficient energy transfer to the latter should occur, and explains why no separate tryptophan emission is observable (D.C. Prasher et al. Gene 111:229-233 (1992). The two cysteines in GFP, Cys<sup>46</sup> and Cys<sup>70</sup>, are 24 Å apart, too distant to form a disulfide bridge. Cys<sup>70</sup> is buried, but Cys<sup>46</sup> should be relatively accessible to sulfhydryl-specific reagents. Such a reagent, 5,5'-dithiobis(2-nitrobenzoic acid), is reported to label GFP and quench its fluorescence (S. Inouye & F.I. Tsuji FEBS Lett. 351:211-214 (1994)). This effect was attributed to the necessity for a free sulfhydryl, but could also reflect specific quenching by the 5-thio-2-nitrobenzoate moiety that would be attached to Cys<sup>41</sup>.

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Although the electron density map is for the most part consistent with the proposed structure of the chromophore (D.C. Prasher et al. Gene 111:229-233 (1992); C. W. Cody et al. Biochemistry 32:1212-1218 (1993)) in the cis [Z-] configuration, with no evidence for any substantial fraction of the opposite isomer around the chromophore double bond, difference features are found at >4 [] in the final (F<sub>o</sub>-F<sub>o</sub>) electron density map that can be interpreted to represent either the intact, uncyclized polypeptide or a carbinolamine (inset to Fig. 2C). This suggests that a significant fraction, perhaps as much as 30% of the molecules in the crystal, have failed to undergo the final dehydration reaction. Confirmation of incomplete dehydration comes from electrospray mass spectrometry, which consistently shows that the average masses of both wild-type and S65T GFP (31.086±4 and 31,099.5±4 Da, respectively) are 6-7 Da higher than predicted (31,079 and 31,093 Da, respectively) for the fully matured proteins. Such a discrepancy could be explained by a 30-35% mole fraction of apoprotein or carbinolamine with 18 or 20 Da higher molecular weight The natural abundance of 13C and 2H and the finite resolution of the Hewlett-Packard 5989B electrospray mass spectrometer used to make these measurements do not permit the individual peaks to be resolved, but instead yields an average mass peak with a full width at half maximum of approximately 15 Da. The molecular weights shown include the His-tag,

which has the sequence MRGSHHHHHHH GMASMTGGQQM GRDLYDDDDK DPPAEF (SEQ ID NO:5). Mutants of GFP that increase the efficiency of fluorophore maturation might yield somewhat brighter preparations. In a model for the apoprotein, the Thr<sup>65</sup>-Tyr<sup>66</sup> peptide bond is approximately in the □-helical conformation, while the peptide of Tyr<sup>66</sup>-Gly<sup>67</sup> appears to be tipped almost perpendicular to the helix axis by its interaction with Arg<sup>66</sup>. This further supports the speculation that Arg<sup>66</sup> is important in generating the conformation required for cyclization, and possibly also for promoting the attack of Gly<sup>67</sup> on the carbonyl carbon of Thr<sup>65</sup> (A. B. Cubitt et al. *Trends Biochem. Sci.* 20:448-455 (1995)).

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The results of previous random mutagenesis have implicated several amino acid side chains to have substantial effects on the spectra and the atomic model confirms that these residues are close to the chromophore. The mutations T203I and E222G have profound but opposite consequences on the absorption spectrum (T. Ehrig et al. FEBS Letters 367:163-166 (1995)). T203I (with wild-type Ser<sup>65</sup>) lacks the 475 nm absorbance neak usually attributed to the anionic chromophore and shows only the 395 nm peak thought to reflect the neutral chromophore (R. Heim et al. Proc. Natl. Acad. Sci. USA 91:12501-12504 (1994); T. Ehrig et al. FEBS Letters 367:163-166 (1995)). Indeed. Thr203 is hydrogen-bonded to the phenolic oxygen of the chromophore, so replacement by Ile should hinder ionization of the phenolic oxygen. Mutation of Glu<sup>222</sup> to Gly (T. Ehrig et al. FEBS Letters 367:163-166 (1995)) has much the same spectroscopic effect as replacing Ser<sup>65</sup> by Gly. Ala. Cys. Val. or Thr. namely to suppress the 395 nm peak in favor of a peak at 470-490 nm (R. Heim et al. Nature 373:664-665 (1995); S. Delagrave et al. Bio/Technology 13:151-154 (1995)). Indeed Glu<sup>222</sup> and the remnant of Thr<sup>65</sup> are hydrogen-bonded to each other in the present structure, probably with the uncharged carboxyl of Glu<sup>222</sup> acting as donor to the side chain oxygen of Thr65. Mutations E222G, S65G, S65A, and S65V would all suppress such H-bonding. To explain why only wild-type protein has both excitation peaks, Ser65, unlike Thr65, may adopt a conformation in which its hydroxyl donates a hydrogen bond to and stabilizes Glu222 as an anion, whose charge then inhibits ionization of the chromophore. The structure also explains why some mutations seem neutral. For example. Gln<sup>80</sup> is a surface residue far removed from the chromophore, which explains why its accidental and ubiquitous mutation to Arg seems to have no obvious intramolecular spectroscopic effect (M. Chalfie et al. Science 263:802-805 (1994)).

The development of GFP mutants with red-shifted excitation and emission

maxima is an interesting challenge in protein engineering (A. B. Cubitt et al. Trends Biochem. Sci. 20:448-455 (1995); R. Heim et al. Nature 373:664-665 (1995); S. Delagrave et al. Bio/Technology 13:151-154 (1995)). Such mutants would also be valuable for avoidance of cellular autofluorescence at short wavelengths, for simultaneous multicolor reporting of the activity of two or more cellular processes, and for exploitation of fluorescence resonance energy transfer as a signal of protein-protein interaction (R. Heim & R. Y. Tsien, Current Biol. 6:178-182 (1996)). Extensive attempts using random mutagenesis have shifted the emission maximum by at most 6 nm to longer wavelengths, to 514 nm (R. Heim & R.Y. Tsien. Current Biol. 6:178-182 (1996)); previously described "red-shifted" mutants merely suppressed the 395 nm excitation peak in favor of the 475 nm 10 peak without any significant reddening of the 505 nm emission (S. Delagrave et al. Bio/Technology 13:151-154 (1995)). Because Thr<sup>203</sup> is revealed to be adjacent to the phenolic end of the chromophore, we mutated it to polar aromatic residues such as His, Tyr. and Trp in the hope that the additional polarizability of their 

systems would lower the 15 energy of the excited state of the adjacent chromophore. All three substitutions did indeed shift the emission peak to greater than 520 nm (Table F). A particularly attractive mutation was T203Y/S65G/V68L/S72A, with excitation and emission peaks at 513 and 527 nm respectively. These wavelengths are sufficiently different from previous GFP mutants to be readily distinguishable by appropriate filter sets on a fluorescence microscope. The 20 extinction coefficient, 36,500 M<sup>-1</sup>cm<sup>-1</sup>, and quantum yield, 0.63, are almost as high as those of S65T (R. Heim et al. Nature 373:664-665 (1995)).

Comparison of Aequorea GFP with other protein pigments is instructive.

Unfortunately, its closest characterized homolog, the GFP from the sea pansy Renilla reniformis (O. Shimomura and F.H. Johnson J. Cell. Comp. Physiol. 59:223 (1962); J. G.

Morin and J. W. Hastings, J. Cell. Physiol. 77:313 (1971); H. Morise et al. Biochemistry 13:2656 (1974); W. W. Ward Photochem. Photobiol. Reviews (Smith, K. C. ed.) 4:1 (1979); W. W. Ward. Bioluminescence and Chemiluminescence (M. A. DeLuca and W. D. McElroy, eds) Academic Press pp. 235-242 (1981); W. W. Ward & S. H. Bokman Biochemistry 21:4535-4540 (1982); W. W. Ward et al. Photochem. Photobiol. 35:803-808 (1982)), has not been sequenced or cloned, though its chromophore is derived from the same FSYG sequence as in wild-type Aequorea GFP (R. M. San Pietro et al. Photochem. Photobiol. 57:63S (1993)). The closest analog for which a three dimensional structure is

available is the photoactive yellow protein (PYP, G. E. O. Borgstahl et al. Biochemistry 34:6278-6287 (1995)), a 14-kDa photoreceptor from halophilic bacteria. PYP in its native dark state absorbs maximally at 446 nm and transduces light with a quantum yield of 0.64, rather closely matching wild-type GFP's long wavelength absorbance maximum near 475 nm and fluorescence quantum yield of 0.72-0.85. The fundamental chromophore in both proteins is an anionic p-hydroxycinnamyl group, which is covalently attached to the protein via a thioester linkage in PYP and a heterocyclic iminolactam in GFP. Both proteins stabilize the negative charge on the chromophore with the help of buried cationic arginine and neutral glutamic acid groups, Arg<sup>52</sup> and Glu<sup>46</sup> in PYP and Arg<sup>96</sup> and Glu<sup>222</sup> in GFP. though in PYP the residues are close to the oxyohenyl ring whereas in GFP they are nearer the carbonyl end of the chromophore. However, PYP has an overall \$\Boxed{\Boxes}\$ fold with appropriate flexibility and signal transduction domains to enable it to mediate the cellular phototactic response, whereas GFP is a much more regular and rigid \( \subseteq \text{-barrel to minimize} \) parasitic dissipation of the excited state energy as thermal or conformational motions. GFP is an elegant example of how a visually appealing and extremely useful function, efficient fluorescence, can be spontaneously generated from a cohesive and economical protein structure.

## A. Summary Of GFP Structure Determination

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Data were collected at room temperature in house using either Molecular 20 Structure Corp. R-axis II or San Diego Multiwire Systems (SDMS) detectors (Cu K II) and later at beamline X4A at the Brookhaven National Laboratory at the selenium absorption edge ((1 = 0.979 Å) using image plates. Data were evaluated using the HKL package (Z. Otwinowski, in Proceedings of the CCP4 Study Weekend: Data Collection and Processing, L. Sawyer, N. Issacs, S. Bailey, Eds. (Science and Engineering Research Council (SERC), 25 Daresbury Laboratory, Warrington, UK, (1991)), pp 56-62; W. Minor, XDISPLAYF (Purdue University, West Lafavette, IN, 1993)) or the SDMS software (A. J. Howard et al. Meth. Enzymol. 114:452-471 (1985)). Each data set was collected from a single crystal. Heavy atom soaks were 2 mM in mother liquor for 2 days. Initial electron density maps were based on three heavy atom derivatives using in-house data, then later were replaced 30 with the synchrotron data. The EMTS difference Patterson map was solved by inspection, then used to calculate difference Fourier maps of the other derivatives. Lack of closure

refinement of the heavy atom parameters was performed using the Protein package (W. Steigemann, in Ph.D. Thesis (Technical University, Munich, 1974)). The MIR maps were much poorer than the overall figure of merit would suggest, and it was clear that the EMTS isomorphous differences dominated the phasing. The enhanced anomalous occupancy for the synchrotron data provided a partial solution to the problem. Note that the phasing power was reduced for the synchrotron data, but the figure of merit was unchanged. All experimental electron density maps were improved by solvent flattening using the program DM of the CCP4 (CCP4: A Suite of Programs for Protein Crystallography (SERC Daresbury Laboratory, Warrington WA4 4AD UK, 1979)) package assuming a solvent content of 38%. Phase combination was performed with PHASCO2 of the Protein package using a weight of 1.0 on the atomic model. Heavy atom parameters were subsequently improved by refinement against combined phases. Model building proceeded with FRODO and O (T. A. Jones et al. Acta, Crystallogr, Sect. A 47:110 (1991); T. A. Jones, in Computational Crystallography D. Savre, Ed. (Oxford University Press, Oxford, 1982) pp. 1.5 303-317) and crystallographic refinement was performed with the TNT package (D. E. Tronrud et al. Acta Cryst. A 43:489-503 (1987)). Bond lengths and angles for the chromophore were estimated using CHEM3D (Cambridge Scientific Computing). Final refinement and model building was performed against the X4A selenomethione data set, using (2F<sub>o</sub>-F<sub>c</sub>) electron density maps. The data beyond 1.9 Å resolution have not been used at this stage. The final model contains residues 2-229 as the terminal residues are not visible in the electron density map, and the side chains of several disordered surface residues have been omitted. Density is weak for residues 156-158 and coordinates for these residues are unreliable. This disordering is consistent with previous analyses showing that residues 1 and 233-238 are dispensible but that further truncations may prevent fluorescence (J. Dopf & T.M. Horiagon. Gene 173:39-43 (1996)). The atomic model has been deposited 25 in the Protein Data Bank (access code 1EMA).

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Table E

## Diffraction Data Statistics

Crystal	Resoluti on (Å)	Total obs	Unique obs	Compl. (%)*	Compl. (shell) <sup>b</sup>	Rmerge (%)°	Riso (%)d
R-axix II							
Native	2.0	51907	13582	80	69	4.1	5.8
EMTS*	2.6	17727	6787	87	87	5.7	20.6
S <b>eM</b> et	2.3	44975	10292	92	88	10.2	9.3
Multiwire	<u> </u>						
HGI4-Se	3.0	15380	4332	84	79	7.2	28.8
<u>X4a</u>							
S <b>eM</b> et	1.8	126078	19503	80	55	9.3	9.4
EMTS	2.3	57812	9204	82	66	7.2	26.3

## Phasing Statistics

Derivative	Resolution (Å)	Number of sites	Phasing power <sup>f</sup>	Phasing Power(shell)	FOM <sup>8</sup>	FOM (shell)
In House						
EMTS	3.0	2	2.08	2.08	0.77	.072
SeMet	3.0	4	1.66	1.28	-	-
HGI4-Se	3.0	9	1.77	1.90	-	-
X4a						
EMTS	3.0	2	1.36	1.26	0.77	.072
SeMet	3.0	4	1.31	1.08	-	-

Atomic Model Statistics Protein atoms 1790 Solvent atoms 94 Resol. range (Å) 20-1.9 Number of reflections (F > 0) 17676 Completeness 84. R. factor(h) 0.175 Mean B-value (Å2) 24.1 10 Deviations from ideality Bond lengths (Å) 0.014 Bond angles (□) 1.9 Restrained B-values (Å2) 4.3 Ramachandran outliers 15 0

Notes:

- (a) Completeness is the ratio of observed reflections to theoretically possible expressed as a percentage.
- (b) Shell indicates the highest resolution shell, typically 0.1-0.4 Å wide.
- (c) Rmerge = □ |I <|>| / □ I, where <|> is the mean of individual observations of intensities I.
- (d) Riso =  $\square |I_{DER} I_{NAT}| / \square I_{NAT}$

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- (e) Derivatives were EMTS=ethymercurithiosalicylate (residues modified Cys<sup>49</sup> and Cys<sup>29</sup>), SeMet=selenomethionine substituted protein (Met¹ and Met³<sup>33</sup> could not be located); Hgl<sub>4</sub>-SeMet = double derivative Hgl<sub>4</sub> on SeMet background.
- 10 (f) Phasing power = <F<sub>H</sub>>/<E> where <F<sub>H</sub>>=r.m.s. heavy atom scattering and <E>=lack of closure.
  - (g) FOM, mean figure of merit
  - (h) Standard crystallographic R-factor, R = □ ||F<sub>obs</sub>| |F<sub>calc</sub>|| / □ |F<sub>obs</sub>|

# 15 B. Spectral properties of Thr<sup>203</sup> ("T203") mutants compared to S65T

The mutations F64L, V68L and S72A improve the folding of GFP at  $37\,\Box$ 

(B. P. Cormack et al. *Gene* 173:33 (1996)) but do not significantly shift the emission spectra.

TABLE F

Clone	Mutations	Excitation max.(nm)	Extinction coefficient (10³M <sup>-1</sup> cm <sup>-1</sup> )	Emission max.(nm)
S65T	S65T	489	39.2	511
5B	T203H/S65T	512	19.4	524
6C	T203Y/S65T	513	14.5	525
10B	T203Y/F64L/S65G/S72A	513	30.8	525
10C	T203Y/F65G/V68L/S72A	513	36.5	527
11	T203W/S65G/S72A	502	33.0	512

WO 98/06	<b>737</b> 51			PCT/US97/14593
12H	T203Y/S65G/S72A	513	36.5	527
20A	T203Y/S65G/V68L/Q69K/S72A	515	46.0	527

The present invention provides novel long wavelength engineered fluorescent proteins. While specific examples have been provided, the above description is illustrative and not restrictive. Many variations of the invention will become apparent to those skilled in the art upon review of this specification. The scope of the invention should, therefore, be determined not with reference to the above description, but instead should be determined with reference to the appended claims along with their full scope of equivalents.

All publications and patent documents cited in this application are

10 incorporated by reference in their entirety for all purposes to the same extent as if each individual publication or patent document were so individually denoted.

#### SEQUENCE LISTING

#### (1) GENERAL INFORMATION:

- (i) APPLICANT: The Regents of the University of California et al.
- (ii) TITLE OF INVENTION: LONG WAVELENGTH MUTANT FLUORESCENT PROTEINS
- (iii) NUMBER OF SEQUENCES: 4

#### (iv) CORRESPONDENCE ADDRESS:

- (A) ADDRESSEE: Fish & Richardson P.C.
  - (B) STREET: 4225 Executive Square, Suite 1400
  - (C) CITY: La Jolla
  - (D) STATE: CA
  - (E) COUNTRY: USA (F) ZIP: 92037
- (v) COMPUTER READABLE FORM:
  - (A) MEDIUM TYPE: Floppy disk
  - (B) COMPUTER: IBM PC compatible (C) OPERATING SYSTEM: PC-DOS/MS-DOS
  - (D) SOFTWARE: PatentIn Release #1.0, Version #1.30
- (vii) PRIOR APPLICATION DATA:
  (A) APPLICATION NUMBER: US 60/024,050
  - (B) FILING DATE: 16-AUG-1996 (C) CLASSIFICATION:
- (vii) PRIOR APPLICATION DATA: (A) APPLICATION NUMBER: US 08/706,408
  - (B) FILING DATE: 30-AUG-1996
  - (C) CLASSIFICATION:
- (viii) ATTORNEY/AGENT INFORMATION:
  - (A) NAME: Haile, Lisa A. (B) REGISTRATION NUMBER: 38,347
  - (C) REFERENCE/DOCKET NUMBER: 07257/056W01
  - (ix) TELECOMMUNICATION INFORMATION:
    - (A) TELEPHONE: 619/678-5070 (B) TELEFAX: 619/678-5099
- (2) INFORMATION FOR SEQ ID NO:1:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 716 base pairs (B) TYPE: nucleic acid

    - (C) STRANDEDNESS: single (D) TOPOLOGY: linear
  - (ix) FEATURE:
    - (A) NAME/KEY: CDS (B) LOCATION: 1..714

(xi	) SE	QUEN	CE D	ESCR:	IPTIC	ON: S	SEQ :	ID N	0:1:						
ATG AGT Met Ser 1															48
GAA TTA Glu Leu															96
GGT GAF															144
ACT ACT Thr Thr	Gly														192
TCT TAT Ser Tyr 65															240
CAT GAC His Asp															288
ACT ATA															336
AAG TTT Lys Phe															384
GAT TTT Asp Phe 130	Lys														432
TAT AAC Tyr Asn 145															480
ATC AAA Ile Lys															528
CAA CTA Gln Leu	GCA Ala	GAC Asp 180	TAT Tyr	TAT Tyr	CAA Gln	CAA Gln	AAT Asn 185	ACT Thr	CCA Pro	ATT Ile	CTC Leu	GAT Asp 190	GGC Gly	CCT Pro	576
GTC CTT Val Lev															624
AAA GAT Lys Asp 210	Pro														672
ACA GCT Thr Ala 225															714
TA															716

## (2) INFORMATION FOR SEQ ID NO:2:

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 238 amino acids
(B) TYPE: amino acid

#### (D) TOPOLOGY: linear

#### (ii) MOLECULE TYPE: protein

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:2:

Met Ser Lys Gly Glu Glu Leu Phe Thr Ala Val Val Pro Ile Leu Val

Glu Leu Asp Gly Asp Val Asn Gly His Lys Phe Ser Val Ser Gly Glu 20 25 30

Gly Glu Gly Asp Val Thr Tyr Gly Lys Leu Thr Leu Lys Phe Ile Cys  $_{35}$   $_{40}$ 

Thr Thr Gly Lys Leu Pro Val Pro Trp Pro Thr Leu Val Thr Thr Phe

Ser Tyr Gly Val Gln Cys Phe Ser Arg Tyr Pro Asp His Met Lys Arg 65 70 75 80

His Asp Phe Phe Lys Ser Ala Met Pro Glu Gly Tyr Val Gln Gln Arg 85 90 95

Thr Ile Phe Phe Lys Asp Asp Gly Asn Tyr Lys Thr Arg Ala Glu Val

Lys Phe Glu Gly Asp Thr Leu Val Asn Arg Ile Glu Leu Lys Gly Ile

Asp Phe Lys Glu Asp Gly Asn Ile Leu Gly His Lys Leu Glu Tyr Asn 130 140

Tyr Asn Ser His Asn Val Tyr Ile Met Ala Asp Lys Gln Lys Asn Gly 145 150 155 160

Ile Lys Val Asn Phe Lys Ile Arg His Asn Ile Glu Asp Gly Ser Val

Gln Leu Ala Asp Tyr Tyr Gln Gln Asn Thr Pro Ile Leu Asp Gly Pro
180 185 190

Val Leu Leu Pro Asp Asn His Tyr Leu Ser Thr Gln Ser Ala Leu Ser

Lys Asp Pro Asn Glu Lys Arg Asp His Met Val Leu Leu Glu Phe Val 210 220 Thr Ala Ala Gly Ile Thr His Gly Met Asp Glu Leu Tyr Lys 225 230 235

(2) INFORMATION FOR SEQ ID NO:3:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 720 base pairs
    - (B) TYPE: nucleic acid
    - (C) STRANDEDNESS: single (D) TOPOLOGY: linear
- (ix) FEATURE:

  - (A) NAME/KEY: CDS (B) LOCATION: 1..720

	(xi)	SEC	UEN	CE DE	ESCR	PTIC	ON: S	SEQ :	DN	3:3:						
ATG Met	GTG Val 240	AGC Ser	AAG Lys	GGC Gly	GAG Glu	GAG Glu 245	CTG Leu	TTC Phe	ACC Thr	GGG Gly	GTG Val 250	GTG Val	CCC Pro	ATC Ile	CTG Leu	48
GTC Val 255		CTG Leu														96
GAG Glu																144
		ACC Thr														192
TTC Phe		TAC Tyr 305														240
CAG Gln																288
CGC Arg 335																336
G <b>T</b> G Val																384
ATC Ile																432
AAC Asn	TAC Tyr	AAC Asn 385	AGC Ser	CAC His	AAC Asn	GTC Val	TAT Tyr 390	ATC Ile	ATG Met	GCC Ala	GAC Asp	AAG Lys 395	CAG Gln	AAG Lys	AAC Asn	480
GGC Gly																528
GTG Val 415																576
CCC																624
AGC Ser																672
GTG Val	ACC Thr	GCC Ala 465	GCC Ala	GGG Gly	ATC Ile	ACT Thr	CAC His 470	GGC Gly	ATG Met	GAC Asp	GAG Glu	CTG Leu 475	TAC Tyr	AAG Lys	AAT *	720

<sup>(2)</sup> INFORMATION FOR SEQ ID NO:4:

<sup>(</sup>i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 240 amino acids
(B) TYPE: amino acid
(D) TOPOLOGY: linear

#### (ii) MOLECULE TYPE: protein

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:4:

 Met 1
 Val Ser Lys Gly Glu Glu Leu Phe Thr Gly Val Val Pro 115 Leu 15

 Val Glu Leu Asp Gly Asp Val Asn Gly His Lys Phe Ger Val Ser Gly 25

 Glu Gly Glu Gly Asp Asp Val Asn Gly His Lys Phe Ser Val Ser Gly 25

 Gly Gly Gly Asp Asp Asp Asp Gly Asp Val Asn Gly His Lys Lys Leu Thr Leu Lys Phe He Ile 45

 Cys Thr Thr Gly Lys Leu Pro Val Gln Cys Phe Ala Arg Tyr Pro Asp His Met Lys 86

 Gln Gln Asp Phe Phe Lys Ser Ala Met Pro Glu Gly Tyr Val Gln Glu Gly 70

 Arg Thr 11e Phe Phe Lys Asp Asp Gly Asn Tyr Lys Thr Arg Ala Glu 110

 Val Lys Phe Glu Gly Asp Thr Leu Val Asn Arg His Lys Lys Gly 115

 11e Asp Phe Lys Asp Asp Gly Asn Tyr Lys Thr Arg Ala Glu 112

 11e Asp Phe Lys Asp Asp Gly Asn Leu Gly His Lys Lys Gly 115

 12e Asp Pro Asp Asp Gly Asn Tyr Leu Gly His Lys Lys Gly 116

 Asp Tyr Asn Ser His Asn Val Tyr Ile Met Ala Asp Lys Glu Lys Asp 166

 Gly Tle Lys Val Asp Phe Lys Ile Arg His Asn The Pro Glu Asp Gly Asp Gly 175

 Pro Val Leu Leu Pro Asp Asn His Tyr Gln Gln Asn Thr Pro Ile Glu Asp Gly 196

 Pro Val Leu Leu Pro Asp Asn His Tyr Leu Ser Tyr Gln Ser Ala Leu 200

 Ser Lys Asp Pro Asn Glu Lys Arg Asp His Met Val Leu Leu Glu Phe 215

 Yaz Thr Ala Ala Gly Ile Thr His Gly Met Asp Glu Leu Tyr Lys 24

## WHAT IS CLAIMED IS:

1	
2	1. A nucleic acid molecule comprising a nucleotide sequence encoding
3	a functional engineered fluorescent protein whose amino acid sequence is substantially
4	identical to the amino acid sequence of Aequorea green fluorescent protein (SEQ ID NO:2)
5	and which differs from SEQ ID NO:2 by at least the substitution T203X, wherein X is an
6	aromatic amino acid selected from H, Y, W or F, said functional engineered fluorescent
7	protein having a different fluorescent property than Aequorea green fluorescent protein.
8	
1	2. The nucleic acid molecule of claim 1 wherein the amino acid
2	sequence further comprises a substitution at \$65, wherein the substitution is selected from
3	S65G, S65T, S65A, S65L, S65C, S65V and S65I.
1	
1	3. The nucleic acid molecule of claim 1 wherein the amino acid
2	sequence differs by no more than the substitutions S65T/T203H; S65T/T203Y;
3	S72A/F64L/S65G/T203Y; S72A/S65G/V68L/T203Y; S65G/V68L/Q69K/S72A/T203Y;
4	S65G/S72A/T203Y; or S65G/S72A/T203W.
1	4. The nucleic acid molecule of claim 1 or 2 wherein the amino acid
2	sequence further comprises a substitution at Y66, wherein the substitution is selected from
3	Y66H, Y66F, and Y66W.
1	5. The nucleic acid molecule of claim 1 or 2 wherein the amino acid
2	sequence further comprises a mutation from Table A.
	•
1	6. The nucleic acid molecule of claim 1 or 2 wherein the amino acid
1	o. The nucleic acid molecule of claim 1 of 2 wherein the amino acid

sequence further comprises a folding mutation.

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1	7. The nucleic acid molecule of any of claims 1-3 wherein the
2	nucleotide sequence encoding the protein differs from the nucleotide sequence of SEQ ID
3	NO:1 by the substitution of at least one codon by a preferred mammalian codon.
1	8. The nucleic acid molecule of any of claims 1-3 encoding a fusion
2	protein wherein the fusion protein comprises a polypeptide of interest and the functional
3	engineered fluorescent protein.
1	An expression vector comprising expression control sequences
2	operatively linked to a nucleic acid molecule comprising a nucleotide sequence encoding a
3	functional engineered fluorescent protein whose amino acid sequence is substantially
4	identical to the amino acid sequence of Aequorea green fluorescent protein (SEQ ID NO:2
5	and which differs from SEQ ID NO:2 by at least the amino acid substitution T203X,
6	wherein X is an aromatic amino acid selected from H, Y, W or F, said functional engineer
7	fluorescent protein having a different fluorescent property than Aequorea green fluorescen
8	protein.
1	10. The expression vector of claim 9 wherein the amino acid sequence
2	further comprises a substitution at S65, wherein the substitution is selected from S65G,
3	S65T, S65A, S65L, S65C, S65V and S65I.
1	11. The expression vector of claim 9 wherein the amino acid sequence
2	differs by no more than the substitutions S65T/T203H; S65T/T203Y;
3	S72A/F64L/S65G/T203Y; S72A/S65G/V68L/T203Y; S65G/V68L/Q69K/S72A/T203Y,
4	S65G/S72A/T203Y; or S65G/S72A/T203W.
1	12. The expression vector of claim 10 or 11 wherein the amino acid
2	sequence further comprises a substitution at Y66, wherein the substitution is selected from
3	Y66H, Y66F, and Y66W.

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1 13. The expression vector of claim 10 or 11 wherein the amino acid sequence further comprises a mutation from Table A.

- The expression vector of claim 9 or 10 wherein the amino acid sequence further comprises a folding mutation.
- 1 15. The expression vector of any of claims 9-11 wherein the nucleotide
  2 sequence encoding the protein differs from the nucleotide sequence of SEQ ID NO:1 by the
  3 substitution of at least one codon by a preferred mammalian codon.
- 1 16. The expression vector of any of claims 9-11 encoding a fusion
  2 protein wherein the fusion protein comprises a polypeptide of interest and the functional
  3 engineered fluorescent protein.
- 17. A recombinant host cell comprising an expression vector that comprises expression control sequences operatively linked to a nucleic acid molecule comprising a nucleotide sequence encoding a functional engineered fluorescent protein whose amino acid sequence is substantially identical to the amino acid sequence of Aequorea green fluorescent protein (SEQ ID NO:2) and which differs from SEQ ID NO:2 by at least the amino acid substitution T203X, wherein X is an aromatic amino acid selected from H, Y, W or F, said functional engineered fluorescent protein having a different fluorescent protein brooperty than Aequorea green fluorescent protein.
- 1 18. The recombinant host cell of claim 17 wherein the amino acid sequence further comprises a substitution at S65, wherein the substitution is selected from S65G, S65T, S65A, S65L, S65C, S65V and S65I.

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	19. The	recombinant host cell of claim 17 wherein the amino acid
2	sequence differs by no mor	e than the substitutions S65T/T203H; S65T/T203Y;
3	S72A/F64L/S65G/T203Y;	S72A/S65G/V68L/T203Y; S65G/V68L/Q69K/S72A/T203Y;
ı	S65G/S72A/T203Y; or S65	6G/\$72A/T203W.
	20. The	recombinant host cell of claim 17 or 18 wherein the amine acid
2	sequence further comprises	a substitution at Y66, wherein the substitution is selected from
3	Y66H, Y66F, and Y66W.	
L	21. The	recombinant host cell of claim 17 or 18 wherein the amino acid
2	sequence further comprises	s a mutation from Table A.
		·
L	22. The	recombinant host cell of claim 17 or 18 wherein the amino acid
2	sequence further comprise	s a folding mutation.
		<b>V</b>
1	23. The	recombinant host cell of any of claims 17-19 wherein the
2	nucleotide sequence encod	ling the protein differs from the nucleotide sequence of SEQ ID
3	NO:1 by the substitution of	f at least one codon by a preferred mammalian codon.
1	24. The	recombinant host cell of any of claims 17-19 encoding a fusion
2	protein wherein the fusion	protein comprises a polypeptide of interest and the functional
3	engineered fluorescent pro	otein.
1	25. The	recombinant host cell of any of claims 17-19 which is a
2	prokaryotic cell.	
1	26. The	e recombinant host cell of any of claims 17-19 which is a
2	eukaryotic cell.	

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fluorescent protein.

1	27. A functional engineered fluorescent protein whose amino acid
2	sequence is substantially identical to the amino acid sequence of Aequorea green fluorescent
3	protein (SEQ ID NO:2) and which differs from SEQ ID NO:2 by at least the amino acid
4	substitution T203X, wherein X is an aromatic amino acid selected from H, Y, W or F, said
5	functional engineered fluorescent protein having a different fluorescent property than
6	Aequorea green fluorescent protein.
1	28. The protein of claim 27 wherein the amino acid sequence further
2	comprises a substitution at S65, wherein the substitution is selected from S65G, S65T,
3	S65A, S65L, S65C, S65V and S65I.
1	29. The protein of claim 27 wherein the amino acid sequence differs by
2	no more than the substitutions S65T/T203H; S65T/T203Y; S72A/F64L/S65G/T203Y;
3	\$72A/\$65G/V68L/T203Y; \$65G/V68L/Q69K/\$72A/T203Y; \$65G/\$72A/T203Y; or
4	S65G/ <b>S72</b> A/T203W.
1	30. The protein of claim 27 or 28 wherein the amino acid sequence
2	further comprises a substitution at Y66, wherein the substitution is selected from Y66H,
3	Y66F, and Y66W.
1	31. The protein of claim 27 or 28 wherein the amino acid sequence
2	further comprises a folding mutation.
1	32. The protein of any of claims 27-29 which is a fusion protein wherein
2	the fusion protein comprises a polypeptide of interest and the functional engineered

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1	33. A fluorescently labelled antibody comprising an antibody coupled to
2	a functional engineered fluorescent protein whose amino acid sequence is substantially
3	identical to the amino acid sequence of Aequorea green fluorescent protein (SEQ ID NO:2)
4	and which differs from SEQ ID NO:2 by at least the amino acid substitution T203X,
5	wherein X is an aromatic amino acid selected from H, Y, W or F, said functional engineered
6	fluorescent protein having a different fluorescent property than Aequorea green fluorescent
7	protein.
1	34. The fluorescently labelled antibody of claim 33 wherein the amino
2	acid sequence further comprises a substitution at S65, wherein the substitution is selected
3	from S65G, S65T, S65A, S65L, S65C, S65V and S65I.
1	35. The fluorescently labelled antibody of claim 33 wherein the amino
2	acid sequence differs by no more than the substitutions S65T/T203H; S65T/T203Y;
3	S72A/F64L/S65G/T203Y; S72A/S65G/V68L/T203Y; S65G/V68L/Q69K/S72A/T203Y;
4	S65G/S72A/T203Y; or S65G/S72A/T203W.
1	36. The fluorescently labelled antibody of claim 33 or 34 wherein the
2	amino acid sequence further comprises a substitution at Y66, wherein the substitution is
_	The state of the s

- selected from Y66H, Y66F, and Y66W.

  37. The fluorescently labelled antibody of any of claims 33-35 which is a
- fusion protein wherein the fusion protein comprises the antibody fused to the functional
   engineered fluorescent protein.

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1	38.	A nucleic acid molecu	le comprising a nucleotide sequence encodit	ng				
2	an antibody fused	to a nucleotide sequence e	ncoding a functional engineered fluorescent					
3	protein whose ami	ino acid sequence is substa	ntially identical to the amino acid sequence	of				
4	Aequorea green fl	uorescent protein (SEQ ID	NO:2) and which differs from SEQ ID NO:	2				
5	by at least the ami	no acid substitution T2032	ζ, wherein X is an aromatic amino acid selec	cted				
6	from H, Y, W or I	, said functional engineere	d fluorescent protein having a different					
7	fluorescent proper	rty than Aequorea green flu	orescent protein.					
1	39.	. The nucleic acid mole	cule of claim 38 wherein the amino acid					
2	sequence further of	comprises a substitution at	S65, wherein the substitution is selected from	m				
3	S65G, S65T, S65	A, S65L, S65C, S65V and	S651.					
1	40.	. The nucleic acid mole	cule of claim 38 wherein the amino acid					
2	sequence differs b	y no more than the substitu	ations S65T/T203H; S65T/T203Y;					

- 3 S72A/F64L/S65G/T203Y; S72A/S65G/V68L/T203Y; S65G/V68L/O69K/S72A/T203Y; S65G/S72A/T203Y: or S65G/S72A/T203W.
  - The nucleic acid molecule of claim 38 or 39 wherein the amino acid sequence further comprises a substitution at Y66, wherein the substitution is selected from Y66H, Y66F, and Y66W.
  - 42. A fluorescently labelled nucleic acid probe comprising a nucleic acid probe coupled to a functional engineered fluorescent protein whose amino acid sequence is substantially identical to the amino acid sequence of Aequorea green fluorescent protein (SEQ ID NO:2) and which differs from SEQ ID NO:2 by at least the amino acid substitution T203X, wherein X is an aromatic amino acid selected from H, Y, W or F, said functional engineered fluorescent protein having a different fluorescent property than Aequorea green fluorescent protein.

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The fluorescently labelled nucleic acid probe of claim 42 wherein the amino acid sequence further comprises a substitution at \$65, wherein the substitution is selected from S65G, S65T, S65A, S65L, S65C, S65V and S65L

The fluorescently labelled nucleic acid probe of claim 42 wherein the 44 amino acid sequence differs by no more than the substitutions S65T/T203H; S65T/T203Y; S72A/F64L/S65G/T203Y; S72A/S65G/V68L/T203Y; S65G/V68L/Q69K/S72A/T203Y; S65G/S72A/T203Y; or S65G/S72A/T203W.

45. The nucleic acid molecule of claim 42 or 43 wherein the amino acid sequence further comprises a substitution at Y66, wherein the substitution is selected from Y66H, Y66F, and Y66W.

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46. A nucleic acid molecule comprising a nucleotide sequence encoding a functional engineered fluorescent protein whose amino acid sequence is substantially identical to the amino acid sequence of Aeguorea green fluorescent protein (SEO ID NO:2) and which differs from SEQ ID NO:2 by at least an amino acid substitution at L42, V61, T62, V68, Q69, Q94, N121, Y145, H148, V150, F165, I167, Q183, N185, L220, E222 (not E222G), or V224, said functional engineered fluorescent protein having a different fluorescent property than Aequorea green fluorescent protein.

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47 The nucleic acid molecule of claim 46 wherein the amino acid substitution is: L42X, wherein X is selected from C, F, H, W and Y, V61X, wherein X is selected from F, Y, H and C,

T62X, wherein X is selected from A, V, F, S, D, N, O, Y, H and C.

6 V68X, wherein X is selected from F, Y and H,

O69X, wherein X is selected from K, R, E and G.

Q94X, wherein X is selected from D, E, H, K and N,

9	N121X, wherein X is selected from F, H, W and Y,
10	Y145X, wherein X is selected from W, C, F, L, E, H, K and Q,
11	H148X, wherein X is selected from F, Y, N, K, Q and R,
12	V150X, wherein X is selected from F, Y and H,
13	F165X, wherein X is selected from H, Q, W and Y,
14	I167X, wherein X is selected from F, Y and H,
15	Q183X, wherein X is selected from H, Y, E and K,
16	N185X, wherein X is selected from D, E, H, K and Q,
17	L220X, wherein X is selected from H, N, Q and T,
18	E222X, wherein X is selected from N and Q or
19	V224X, wherein X is selected from H, N, Q, T, F, W and Y.
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1	48. An expression vector comprising expression control sequences
2	operatively linked to a nucleic acid molecule of comprising a nucleotide sequence encoding
3	a functional engineered fluorescent protein whose amino acid sequence is substantially
4	identical to the amino acid sequence of Aequorea green fluorescent protein (SEQ ID NO:2)
5	and which differs from SEQ ID NO:2 by at least an amino acid substitution at L42, V61,
6	T62, V68, Q69, Q94, N121, Y145, H148, V150, F165, I167, Q183, N185, L220, E222 (not
7	E222G), or V224, said functional engineered fluorescent protein having a different
8	fluorescent property than Aequorea green fluorescent protein.
1	49. The expression vector of claim 48 wherein the amino acid
2	substitution is:
3	L42X, wherein X is selected from C, F, H, W and Y,
4	V61X, wherein X is selected from F, Y, H and C,
5	T62X, wherein X is selected from A, V, F, S, D, N, Q, Y, H and C,
6	V68X, wherein X is selected from F, Y and H,
7	Q69X, wherein X is selected from K, R, E and G,
8	Q94X, wherein X is selected from D, E, H, K and N,

9	N121X, wherein X is selected from F, H, W and Y,
10	Y145X, wherein X is selected from W, C, F, L, E, H, K and Q,
11	H148X, wherein X is selected from F, Y, N, K, Q and R,
12	V150X, wherein X is selected from F, Y and H,
13	F165X, wherein X is selected from H, Q, W and Y,
14	I167X, wherein X is selected from F, Y and H,
15	Q183X, wherein X is selected from H, Y, E and K,
16	N185X, wherein X is selected from D, E, H, K and Q,
17	L220X, wherein X is selected from H, N, Q and T,
18	E222X, wherein X is selected from N and Q or
19	V224X, wherein X is selected from H, N, Q, T, F, W and Y.
1	50. A recombinant host cell comprising an expression vector that
2	comprises expression control sequences operatively linked to a nucleic acid molecule
3	comprising a nucleotide sequence encoding a functional engineered fluorescent protein
4	whose amino acid sequence is substantially identical to the amino acid sequence of
5	Aequorea green fluorescent protein (SEQ ID NO:2) and which differs from SEQ ID NO:2
6	by at least an amino acid substitution at L42, V61, T62, V68, Q69, Q94, N121, Y145,
7	H148, V150, F165, I167, Q183, N185, L220, E222 (not E222G), or V224, said functional
8	engineered fluorescent protein having a different fluorescent property than Aequorea green
9	fluorescent protein.
1	51. The recombinant host cell of claim 50 wherein the amino acid
2	substitution is:
3	L42X, wherein X is selected from C, F, H, W and Y,
4	V61X, wherein X is selected from F, Y, H and C,
5	T62X, wherein X is selected from A, V, F, S, D, N, Q, Y, H and C,
6	V68X, wherein X is selected from F, Y and H,
7	Q69X, wherein X is selected from K, R, E and G,

8	Q94X, wherein X is selected from D, E, H, K and N,
9	N121X, wherein X is selected from F, H, W and Y,
10	Y145X, wherein X is selected from W, C, F, L, E, H, K and Q,
11	H148X, wherein X is selected from F, Y, N, K, Q and R,
12	V150X, wherein X is selected from F, Y and H,
13	F165X, wherein X is selected from H, Q, W and Y,
14	I167X, wherein X is selected from F, Y and H,
15	Q183X, wherein X is selected from H, Y, E and K,
16	N185X, wherein X is selected from D, E, H, K and Q,
17	L220X, wherein X is selected from H, N, Q and T,
18	E222X, wherein X is selected from N and Q or
19	V224X, wherein X is selected from H, N, Q, T, F, W and Y.
20	
1	52. A functional engineered fluorescent protein whose amino acid
2	sequence is substantially identical to the amino acid sequence of Aequorea green fluorescent
3	protein (SEQ ID NO:2) and which differs from SEQ ID NO:2 by at least an amino acid
4	substitution at L42, V61, T62, V68, Q69, Q94, N121, Y145, H148, V150, F165, I167,
5	Q183, N185, L220, E222 (E222G), or V224, said functional engineered fluorescent protein
6	having a different fluorescent property than Aequorea green fluorescent protein.
1	53. The functional engineered fluorescent protein of claim 52 wherein the
2	amino acid substitution is:
3	L42X, wherein X is selected from C, F, H, W and Y,
4	V61X, wherein X is selected from F, Y, H and C,
5 ·	T62X, wherein X is selected from A, V, F, S, D, N, Q, Y, H and C,
6	V68X, wherein X is selected from F, Y and H,
7	Q69X, whèrein X is selected from K, R, E and G,
8	Q94X, wherein X is selected from D, E, H, K and N,

9	N121X, wherein X is selected from F, H, W and Y,
10	Y145X, wherein X is selected from W, C, F, L, E, H, K and Q,
11	H148X, wherein X is selected from F, Y, N, K, Q and R,
12	V150X, wherein X is selected from F, Y and H,
13	F165X, wherein X is selected from H, Q, W and Y,
14	I167X, wherein X is selected from F, Y and H,
15	Q183X, wherein X is selected from H, Y, E and K,
16	N185X, wherein X is selected from D, E, H, K and Q,
17	L220X, wherein X is selected from H, N, Q and T,
18	E222X, wherein X is selected from N and Q or
19	V224X, wherein X is selected from H, N, Q, T, F, W and Y.
1	54. A fluorescently labelled antibody comprising an antibody coupled to
2	a functional engineered fluorescent protein whose amino acid sequence is substantially
3	identical to the amino acid sequence of Aequorea green fluorescent protein (SEQ ID NO:2
4	and which differs from SEQ ID NO:2 by at least an amino acid substitution at L42, V61,
5	T62, V68, Q69, Q94, N121, Y145, H148, V150, F165, I167, Q183, N185, L220, E222 (no
6	E222G), or V224, said functional engineered fluorescent protein having a different
7	fluorescent property than Aequorea green fluorescent protein.
1	55. The antibody of claim 54 wherein the amino acid substitution is:
2	L42X, wherein X is selected from C, F, H, W and Y,
3	V61X, wherein X is selected from F, Y, H and C,
4	T62X, wherein X is selected from A, V, F, S, D, N, Q, Y, H and C,
5	V68X, wherein X is selected from F, Y and H,
6	Q69X, wherein X is selected from K, R, E and G,
7	Q94X, wherein X is selected from D, E, H, K and N,
8	N121X, wherein X is selected from F, H, W and Y,

9	Y145X, wherein X is selected from W, C, F, L, E, H, K and Q,
10	H148X, wherein X is selected from F, Y, N, K, Q and R,
11	V150X, wherein X is selected from F, Y and H,
12	F165X, wherein X is selected from H, Q, W and Y,
13	I167X, wherein X is selected from F, Y and H,
14	Q183X, wherein X is selected from H, Y, E and K,
15	N185X, wherein X is selected from D, E, H, K and Q,
16	L220X, wherein X is selected from H, N, Q and T,
17	E222X, wherein X is selected from N and Q or
18	V224X, wherein X is selected from H, N, Q, T, F, W and Y.
1	56. A nucleic acid molecule comprising a nucleotide sequence encoding
2	an antibody fused to a nucleotide sequence encoding a functional engineered fluorescent
3	protein whose amino acid sequence is substantially identical to the amino acid sequence of
4	Aequorea green fluorescent protein (SEQ ID NO:2) and which differs from SEQ ID NO:2
5	by at least an amino acid substitution at L42, V61, T62, V68, Q69, Q94, N121, Y145,
6	H148, V150, F165, I167, Q183, N185, L220, E222 (not E222G), or V224, said functional
7	engineered fluorescent protein having a different fluorescent property than Aequorea green
8	fluorescent protein.
1	57. The nucleic acid molecule of claim 56 wherein the amino acid
2	substitution is:
3	L42X, wherein X is selected from C, F, H, W and Y,
4	V61X, wherein X is selected from F, Y, H and C,
5	T62X, wherein X is selected from A, V, F, S, D, N, Q, Y, H and C,
6	V68X, wherein X is selected from F, Y and H,
7	Q69X, wherein X is selected from K, R, E and G,
8	Q94X, wherein X is selected from D, E, H, K and N,
9	N121X, wherein X is selected from F, H, W and Y.

10	Y145X, wherein X is selected from W, C, F, L, E, H, K and Q,
11	H148X, wherein X is selected from F, Y, N, K, Q and R,
12	V150X, wherein X is selected from F, Y and H,
13	F165X, wherein X is selected from H, Q, W and Y,
14	I167X, wherein X is selected from F, Y and H,
15	Q183X, wherein X is selected from H, Y, E and K,
16	N185X, wherein X is selected from D, E, H, K and Q,
17	L220X, wherein X is selected from H, N, Q and T,
18	E222X, wherein X is selected from N and Q or
19	V224X, wherein X is selected from H, N, Q, T, F, W and Y.
1	58. A fluorescently labelled nucleic acid probe comprising a nucleic aci
2	probe coupled to a functional engineered fluorescent protein whose amino acid sequence i
3	substantially identical to the amino acid sequence of Aequorea green fluorescent protein
4	(SEQ ID NO:2) and which differs from SEQ ID NO:2 by at least an amino acid substitution
5	at L42, V61, T62, V68, Q69, Q94, N121, Y145, H148, V150, F165, I167, Q183, N185,
6	L220, E222 (E222G), or V224, said functional engineered fluorescent protein having a
7	different fluorescent property than Aequorea green fluorescent protein.
1	59. The probe of claim 58 wherein the amino acid substitution is:
2	L42X, wherein X is selected from C, F, H, W and Y,
3	V61X, wherein X is selected from F, Y, H and C,
4	T62X, wherein X is selected from A, V, F, S, D, N, Q, Y, H and C,
5	V68X, wherein X is selected from F, Y and H,
6	Q69X, wherein X is selected from K, R, E and G,
7	Q94X, wherein X is selected from D, E, H, K and N,
8	N121X, wherein X is selected from F, H, W and Y,
9	Y145X, wherein X is selected from W, C, F, L, E, H, K and Q,

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10	H148X, wherein X is selected from F, Y, N, K, Q and R,
11	V150X, wherein X is selected from F, Y and H,
12	F165X, wherein X is selected from H, Q, W and Y,
13	I167X, wherein X is selected from F, Y and H,
14	Q183X, wherein X is selected from H, Y, E and K,
15	N185X, wherein X is selected from D, E, H, K and Q,
16	L220X, wherein X is selected from H, N, Q and T,
17	E222X, wherein X is selected from N and Q or
18	V224X, wherein X is selected from H, N, Q, T, F, W and Y.
1	60. A method for determining whether a mixture contains a target
2	comprising:
3	contacting the mixture with a fluorescently labelled probe comprising
4	a probe and a functional engineered fluorescent protein of claim 27 or claim 52; and
5	determining whether the target has bound to the probe.
1	61. The method of any of claim 60 the target is bound to a solid matrix.
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2	62. A method for engineering a functional engineered fluorescent protein
3	having a fluorescent property different than Aequorea green fluorescent protein, comprising
4	substituting an amino acid that is located no more than 0.5 nm from any atom in the
5	chromophore of an Aequorea-related green fluorescent protein with another amino acid;
6	whereby the substitution alters a fluorescent property of the protein.
1	63. The method of claim 62 wherein the amino acid substitution alters the
2	electronic environment of the chromophore.
3	

L	64.	A method for engineering a functional engineered fluorescent protein
2	having a different f	luorescent property than Aequorea green fluorescent protein comprising
3	substituting amino	acids in a loop domain of an Aequorea-related green fluorescent protein
1	with amino acids so	as to create a consensus sequence for phosphorylation or for
5	proteolysis.	
L	65.	A method for producing fluorescence resonance energy transfer
2	comprising:	
3		providing a donor molecule comprising a functional engineered
1	fluorescent protein	of claim 27 or claim 52;
5		providing an appropriate acceptor molecule for the fluorescent
5	protein; and	
7		bringing the donor molecule and the acceptor molecule into
3	sufficiently close c	ontact to allow fluorescence resonance energy transfer.
1	66.	A method for producing fluorescence resonance energy transfer
2	comprising:	
3		providing an acceptor molecule comprising a functional engineered
4	fluorescent protein	of claim 27 or claim 52;
5		providing an appropriate donor molecule for the fluorescent protein;
6	and	
7		bringing the donor molecule and the acceptor molecule into
8	sufficiently close of	contact to allow fluorescence resonance energy transfer.
1	67.	The method of claim 66 wherein the donor molecule is a engineered
2	fluorescent proteir	whose amino acid sequence comprises the substitution T203I and the
3	acceptor molecule	is a nutant fluorescent protein whose amino acid sequence comprises the
4	substitution T2032	X, wherein X is an aromatic amino acid selected from H, Y, W or F, said
5	functional enginee	ered fluorescent protein having a different fluorescent property than

Aequorea green fluorescent protein.

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68. A nucleic acid molecule comprising a nucleotide sequence encoding
a functional engineered fluorescent protein whose amino acid sequence is substantially
identical to the amino acid sequence of Aequorea green fluorescent protein (SEQ ID NO:2)
and which differs from SEQ ID NO:2 by at least an amino acid substitution located no more
than about 0.5 nm from the chromophore of the engineered fluorescent protein, wherein the
substitution alters the electronic environment of the chromophore, whereby the functional
engineered fluorescent protein has a different fluorescent property than Aequorea green
fluorescent protein

- 69. An expression vector comprising expression control sequences operatively linked to a nucleotide sequence encoding a functional engineered fluorescent protein whose amino acid sequence is substantially identical to the amino acid sequence of Aequorea green fluorescent protein (SEQ ID NO:2) and which differs from SEQ ID NO:2 by at least an amino acid substitution located no more than about 0.5 nm from the chromophore of the engineered fluorescent protein, wherein the substitution alters the electronic environment of the chromophore, whereby the functional engineered fluorescent protein has a different fluorescent property than Aequorea green fluorescent protein.
- 70. A functional engineered fluorescent protein whose amino acid sequence is substantially identical to the amino acid sequence of Aequorea green fluorescent protein (SEQ ID NO:2) and which differs from SEQ ID NO:2 by at least an amino acid substitution located no more than about 0.5 nm from the chromophore of the engineered fluorescent protein, wherein the substitution alters the electronic environment of the chromophore, whereby the functional engineered fluorescent protein has a different fluorescent property than Aequorea green fluorescent protein.
- 71. A crystal of a protein comprising a fluorescent protein with an amino acid sequence substantially identical to SEQ ID NO: 2, wherein said crystal diffracts with at least a 2.0 to 3.0 angstrom resolution.

1	72. The crystal of claim 71, wherein the fluorescent protein has at least
2	200 amino acids, a completeness value of at least 80% and has a crystal stability within
3	0.5% of its unit cell dimensions.
1	73. The crystal of claim 71, wherein the amino acid sequence comprises a
2	substitution at S65, wherein the substitution is selected from S65G, S65T, S65A, S65L,
3	S65C, S65V and S65I.
1	74. The crystal of claim 71, wherein said crystal has the following unit
2	cell dimensions in angstroms: a = 51.8, b= 62.8 and c= 70.7 with a space group of P2 2 2
3	and an □ angle of 90.00□, a □ angle of 90.00□ and a □ angle of 90.00□ and the crystal has
4	a diffraction limit where 90% or greater of the potential reflections can be used to determine
5	the coordinates of the atoms.
1	75. A computational method of designing a fluoresent protein
2	comprising:
3	determining from a three dimensional model of a crystallized
4	fluorescent protein comprising a fluorescent protein with a bound ligand, at least one
5	interacting amino acid of the fluorescent protein that interacts with at least one first
5	chemical moiety of the ligand, and
7	selecting at least one chemical modification of the first chemical
8	moiety to produce a second chemical moiety with a structure to either decrease or increase
9	an interaction between the interacting amino acid and the second chemical moiety compared
10	to the interaction between the interacting amino acid and the first chemical moiety.
1	76. The computational method of claim 75, further comprising generating

the three dimensional model of the crystallized protein comprising a fluorescent protein

with an amino acid sequence substantially identical to SEQ ID NO:2.

ı	77. The computational method of claim 75, wherein the selecting selects
2	the first chemical moiety that interacts with at least one of the amino acids listed in Figs. 5-
3	to 5-28.
I	78. The computational method of claim 75, wherein the chemical
2	modification enhances hydrogen bonding interaction, charge interaction, hydrophobic
3	interaction, Van Der Waals interaction or dipole interaction between the second chemical
4	moiety and the interacting amino acid compared to the first chemical moiety and the
5	interacting amino acid.
l	79. A computational method of modeling the three dimensional structure
2	of a fluorescent protein comprising determining a three dimensional relationship between a
3	least two atoms listed in the atomic coordinates of Figs. 5-1 to 5-28.
1	80. The computational method of claim 79, wherein the determining
2	comprises determining the three dimensional structure of a fluorescent protein with an
3	amino acid sequence at least 80% identical to SEQ ID NO:2.
4	
1	81. The computational method of claim 79, wherein the determining
2	comprises determining the three dimensional structure of a fluorescent protein with an
3	amino acid sequence at least 95% identical to SEQ ID NO:2.
1	82. The computational method of claim 79, wherein the determining
2	comprises determining the three dimensional relationship of at least 1500 atoms listed in
3	Figs. 5-1 to 5-28.

83. A device comprising a storage device and, stored in the device, at

least 10 atomic coordinates selected from the atomic coordinates listed in Figs. 5-1 to 5-28.

- 1 84. The device of claim 83, wherein the storage device is a computer
- 2 readable device that stores code that receives as input the atomic coordinates.

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1	85.	The device of claim 84, wherein computer readable device is a floppy
2	disk or a hard drive.	

- A nucleic acid molecule comprising a nucleotide sequence encoding a functional 3 86
- engineered fluorescent protein whose amino acid sequence is substantially identical to
- the amino acid sequence of Aeguorea green fluorescent protein (SEO ID NO:2) and 5
- which differs from SEQ ID NO:2 by at least a substitution at O69, wherein said 6
- 7 functional engineered fluorescent protein has a different fluorescent property than
- 8 Aequorea green fluorescent protein.
- The nucleic acid molecule of claim 86, wherein said substitution at Q69 is selected G 87.
- 10 from the group of K, R, E and G.
- 11 88 The nucleic acid molecule of claim 86, wherein said amino acid sequence further
- comprises a function mutation at \$65. 12
- 13 89. A nucleic acid molecule comprising a nucleotide sequence encoding a functional
- 14 engineered fluorescent protein whose amino acid sequence is substantially identical to
- the amino acid sequence of Aequorea green fluorescent protein (SEQ ID NO:2) and 15
- 16 which differs from SEQ ID NO:2 by at least a substitution at E222, but not including
- 17 E222G, wherein said functional engineered fluorescent protein has a different
- 18 fluorescent property than Aequorea green fluorescent protein.
- 19 90. The nucleic acid molecule of claim 89, wherein said substitution at E222 is selected
- 20 from the group of N and Q.

25

- 22 The nucleic acid molecule of claim 89, wherein said amino acid sequence further
- 22 comprises a function mutation at F64.
- A nucleic acid molecule comprising a nucleotide sequence encoding a functional 23 92.
- engineered fluorescent protein whose amino acid sequence is substantially identical to 24
- the amino acid sequence of Aequorea green fluorescent protein (SEQ ID NO:2) and
- which differs from SEQ ID NO:2 by at least a substitution at Y145, wherein said 26
- 27 functional engineered fluorescent protein has a different fluorescent property than
- 28 Aequorea green fluorescent protein.
- 29 The nucleic acid molecule of claim 92, wherein said substitution at Y145 is selected 30 from the group of W, C, F, L, E, H, K and O.

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31	94.	The nucleic acid molecule of claim 92, wherein said amino acid sequence further
32	co	mprises a function mutation at Y66.
33	95.	A method of identifying a test chemical, comprising:
34		contacting a test chemical a sample containing a biological entity labeled with a
35	functi	onal, engineered fluorescent protein or a polynucleotide encoding said functional,
36	engine	eered fluorescent protein, and
37		detecting fluorescence of said functional engineered fluorescent protein.
38	96.	The method of claim 95, wherein said fluorescence in the presence of a test
39	ch	emical is greater than in the absence of said test chemical.
40	97.	The method of claim 96, wherein said polynucleotide encoding said functional,
41	en	gineered fluorescent protein is operatively linked to a genomic polynucleotide.
42	98.	The method of claim 95, wherein said functional, engineered fluorescent protein is
43	fu	sed to second functional protein.
44	99.	The method of claim 96, wherein said polynucleotide encoding said functional,
45	er	gineered fluorescent protein is operatively linked to a response element.
46	100.	The method of claim 96, wherein said polynucleotide encoding said functional,
47	er	gineered fluorescent protein is operatively linked to a response element in a

48

mammalian cell.

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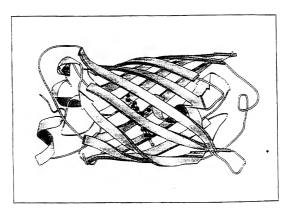


Figure 1a

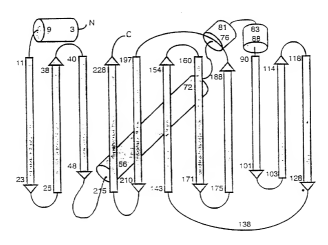


Figure 1b

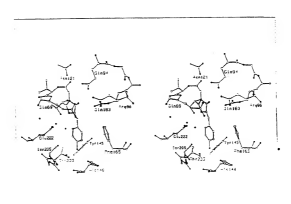


Figure 2a

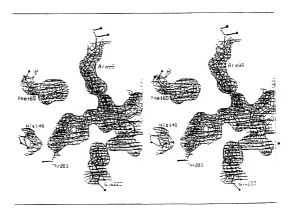


Figure 2b

Figure 2c

## (xi) SEQUENCE DESCRIPTION:

:0:1: :0:2:		Ser				Glu					Val					GT7 Val	-8
					ASD					Lys					GLY	GAS Glu	₹6
	GGT	GAA	607 617 35	ASD	WIS	ACA Thr	TAC	GGA GLY 40	Lys	CTT	ACC	Leu	LYS 45	Phe	lle	750 Cys	144
	ACT	ACT	GGA	***	CTA	ccr	577	CCA	ras	CCA	ACA	CTT	GTC	ACT	ACT	775	192
	The	1h.		Lys	Leu	Pro	val 55		trp	Pro	ihr	60		Thr	Thr	Phe	
	Ser 65	TAT	GLY	GT: Val	GLO	FSC Cys 70	Phe	TCA Ser	AGA	TAC	Pro 75	ASD	HIS	ATG	L ys	CG5 Arg 80	240
	CAT	GAC ASD	777 Phe	TTC Phe	AAG Lys 85	Ser	SSS Ala	ATG	Pro	GLU 90	GGT	TAT	GTA	Gla	61 u 95	AGA	258
		ATA Ite			Lys												236
	AAG Lys	Phe	GAA Glu 115	GGT	CAT Asp	ACC	ctt Leu	GTT Val 120	Asn	AGA Arg	ATC	GAG Glu	77A Leu 125	AAA Lys	GST Gly	ATT [le	384
		777 Phe- 130	Lys														432
		AAC Asn															480
	ile	AAA Lys	GTT Val	AAC Asn	775 Phe 165	LYS	4:1   e	ASA Arg	ELC	AAC Asn 170	ile	GAA	GAT GAT	GGA Gly	AGC Ser 175	STT VAI	125
	Gln	C7A	SC.A Ala	CAC ASD 180	CAT HIS	TAT	Cin Cin	GLA	441 Asn 185	ACT Thr	433 619	ATT [le	GSS Gly	547 ASD 190	GGC Gly	co: Pro	376
	GTC Val	CT7 Leu	774 Leu 195	ECA Pro	GAC ASD	AAC ASD	H.2	TAC 1 yr 200	CTS Leu	TC: Ser	AEA 181	CAA Gln	5 e r 205	GCC Ala	ett Leu	TES Ser	524
	AAA Lys	GA: ASD 210	222 Pro	AAC ASD	GAA GLU	LYS	AGA Arg 215	GAC ASD	EAC #15	ATG Het	Val	211 Leu 220	ett Leu	GAG	777 Phe	STA Val	£72
	ACA Thr 225	730 61A	TEE Ala	GSG GL y	He	ACA Thr 230	r:5	555 51y	AIS Pet	A S D	64A 61u 235	CTA Leu	TAC	AAA Lys	7.4		717

Figure 3

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TOTAL BESS ITTA mumanized codon Usago, Pith an additional amino acid after the start met to provide optimal Kozak Seguence

		:			18			2.7			36			45			54
475	373		443	550	SAS	GAS	C73	772		333	375	575		ATC		375	
445	- 4 -	ier	Lys	Gly	619	Glu	Leu	Pne	inr	Siy	Val	Val	Pro	11=	Leu	al	Giu
		63			72			81			90			99			108
:::5	GAS	220	SAC	GTA	ಭ	SSE	SAS	AA5	TTS	AGC	375	722	035	SAG	GGC	CAG	GSC
Leu	Asp	521.	Asp	Val	Ast	siy	H±s	Lys	Pne	šer	Val	Ser	Gly	Glu	Gly	Siu	Gly
		117			126			135			144			153			162
GAT	900	222	TAC	GGC	AAG	273	ACC	CTS	RAG	775	ATC	755	ACC	ACC	ಯಾ	AAG	CTS
 24D	ALA	755	Tor	Glv	Lva	Leu	The	Leu	Lys	Phe	ile	Cys	The	Thr	Gly	Lvs	Leu
								189			198			207			216
ccc	G73	171	TGG	ccc	180 ACC	cre	STS	ACC	acc	770		TAC			CAG		
									~								
510	Va.	Pro	Trp	Pro	Thr	Leu	Val	Thr	TAT	Phe	Giy	Tyr	GLY	VAI	GIN	Cys	Phe
		225			234			243 CAG			252			261			270
Ala	Arq	77/2	Pro	Asp	илв	Met	Lys	Gln	His	Азр	Pre	Phe	Lys	Ser	Aia	Met	bio
		279			288			297			306			315			324
GAA	GSS	77.5	Gic	CAS	GAG	csc	ACC	ATC	777	777	ZAG.	GAC	GAC	GGC	AAC	TAC	AAG
								11.									
		333			342			351			360			369			378
		300	GAG		AAG	770		ggs	SAC		575	GTS		CGC	ATC		CTG
		41.0	G) 11	Val	Lve	Pne	Glu	Gly	Asp	Thr	Leu	Val	Asn	Aro	Ile	Glu	Leu
														423			
AAG	GGS	387 ATC	GAC	772	396 AAG	GAG	GAC	405 GSC	AAC	ATO	414	GGG	cac	AAG	CT3	GAG	TAC
								Giy									
Lys	G.,		Asp	Prom			vař									0.11	
		441		CAF	450		727	459	a.TG	gee	468		CAG	477		200	486 2TC
Ası		: As:	ser	His	Asr	va3	777	: ::=	Met	A.a	AST	Lys	Gin	LVS	Asn	Gly	11e
		49			50-			513			522			53	L		540
															CAU		
<b>Σ</b> γ	s Va	l As	Phe	Ly	: 11	A.	H1	s Asr	110	Gl:	ASI	Gly	Set	· Va	Gir	Lei	AL.
		54			. 55			567	,		57			58			59
								TA C									
								0 1.4									
		60			61			62:			63			63			64
AA	c ಜ	= TA	c	3 23	TA:	c sa	; TC	r ge		5 45	حد ت	L GAS	: :::	: 22	C SAC	***	3 CG
As			Let										PT:	As	n Glu	Ly	s Ar
	n Hi	s Ty	Len	ı Se	T 7	r G1	. Se	F A1	a Lei	ı Sa	= 1.y	AST		69			70
SA	n Hi T CA	5 TY 55 5 AT	r Len	. se:	66 CT	r Gi 6 3 GA	s se	F A1.	Lei S S AC	: Se : CC	e 57 68 6 90	AST GGG	) AT	69 : AC	3 T CAC	: ga	70:
GA 	n Hi	8 TY 65 8 AT	Let	a Se:	66 CT	r Gl 6 G GA	5 Se	67:	Lei	: Se	e Ly	AST G GGG	ATI	69 . AC	T CAC	: ga	70: AT
GA 	n Hi	8 TY 65 8 AT	T Len	2 Se:	66 CT	r Gl 6 G GA	5 Se	F A1.	Lei	: Se	e Ly	AST G GGG	ATI	69 . AC	T CAC	: ga	70: AT
3A 	T CA	5 TY 65 2 AT  5 Mg	GT:	1 Se:	66 C.	e Gi G GA 	5 Se	67:	Lei	: Se	e Ly	AST G GGG	ATI	69 . AC	T CAC	: ga	70: AT
GA As	T CA	5 TY 65 5 AT 5 Ks	T Len	1 Se:	66 5 CT 4 Le	6 GA G1 G1 G1 G1	5 Se	67:	Lei	: Se	e Ly	AST G GGG	ATI	69 . AC	T CAC	: ga	70: AT

Figure 4

CRYSTI	5.1	767	5.7	845 70.6	566 90.CC	90.00	30.00	
ORIGXI		1.000		0.000000	0.000000		0.00000	
ORIGX2		0.000		1.000000	0.000000		0.00000	
ORIGX3		0.000	0000	0.000000	1.000000	)	0.00000	
SCALE 1		0.019	717	0.000000	0.000000		0.00000	
SCALE 2		0.000		0.015912	0.000000		0.00000	
SCALES		0.000		0.000000	0.014151		0.00000	
ATOM	1	N	SER	2	28.888	9.409	52.301	1.00 85.05
ATOM	2	CA	SER	2	27.638	10.125	52.516	1.00 80.05
ATOM	3	c		2	26.499			1.00 85.35
			SER			9.639	51.644	
MOTA	4	0	SER	2	26.606	8.656	50.915	1.00 84.55
MOTA	5	CB	SER	2	27.783	11.635	52.378	1.00 70.97
MOTA	6	OG	SER	2	27.690	12.033	51.012	1.00 44.08
ATOM	7	N	LYS	3	25.418	10.403	51.731	1.00 87.71
ATOM	é	CA	LYS	š				
					24.141	10.191	E1.036	1.00 87.15
MOTA	9	C	LYS	3	24.214	10.266	49.497	1.00 76.86
MOTA	10	0	LYS	3	24.107	9.258	48.774	1.00 78.27
ATOM	11	CB	LYS	3	23.127	11.240	51.521	1.00 89.44
ATOM	12	CG	LYS	3	21.768	10.697	51.949	1.00 75.06
							21.949	
ATOM	13	CD	LYS	3	20.681	11.781	51.987	1.00 76.53
MOTA	14	CE	LYS	3	20.711	12.655	53.243	1.00 68.55
ATOM	15	NZ	178	3	20.816	14.103	52.953	1.00 46.24
ATOM	16	N	GLY	4	24.318	11.495	49.015	00 53 62
ATOM	17	CA				11.495		1.00 53.62
			GLY	4	24.297	11.798	47.605	1.00 45.97
MOTA	18	C	GLY	4	25.425	11.206	46.796	1.00 31.90 1.00 33.63 1.00 32.54 1.00 32.57
MOTA	19	0	GLY	4	25.234	10.923	45.619	1.00 33.63
ATOM	20	N	GLU	5	26.606	11.082	47.420	1.00 32.54
ATOM	21	CA	GLU	5	27.821	10.598	46.726	1 00 33 57
MOTA	22	c		š	27.523			1.00 32.37
			GLU			9.590	45.616	1.00 28.40
MOTA	23	0	GLU	5	27.850	9.803	44.444	1.00 26.12
ATOM	24	CB	GLU	5	28.873	10.053	47.718	1.00 38.53
MOTA	25	CG	GLU	5	30.337	10.461	47.425	1.00 41.35
ATOM	26	CD	GLU	5	31.311	9.584	48.170	1.00 90.82
							40.170	
ATOM	27		GLU	5	31.508	9.677	49.381	1.00 74.80
ATOM	28		GLU	5	31.839	8.653	47.403	1.00100.00
MOTA	29	N	GLU	á	26.893	3.499	46.017	1.00 23.57
ATOM	30	CA	CLU	6	26.479	7.410	45.150	1.00 31.50
ATOM	31	c	GLU	6	25.551	7.837	43.979	1.00 31.13
	32						43.979	1.00 31.10
ATOM		0	CLU	6	25.479	7.142	42.955	1.00 30.96
HOTA	33	CB	GLU	6	25.780	5.330	45.992	1.00 35.64 1.00 55.53 1.00 22.26
ATOM	34	CG	GLU	6	25.260	5.893	47.338	1.00 55.53
ATOM	35	N	TEA TEA	7	24.864	2.956	44.138	. 00 22 26
ATOM	36	CA		7	23.954	9.456	-3.089	1.00 21.61
	37			ź				
MOTA		C			24.693	10.061	41.917	1.00 16.90
MOTA	38	0	LEU	7	24.152	10.250	÷0.836	1.00 18.38
MOTA	39	CB	LEU	7	23.050	10.548	43.665	1,00 22.41
MOTA	40	CG	LEU	7	21.672	10.058	44.098	1.00 18.38 1.00 22.41 1.00 32.84
ATOM	41	CD1		ż	21.597	8.536	44.074	1.00 31.64
ATOM	42		120	ź	21.597			
			LEU		21.332	10.591	45.485	1.00 33.14
MOTA	43	:1	PHE	8	25.944	10.407	42.157	1.00 20.75
ATOM	44	CA	PHE	9	26.740	11.132	41.159	1.00 20.75
ATOM	45	С	PHE	8	27.818	10.333	-0.427	1.00 30.59
MOTA	46	ò	PHE	8	28.590	10.856	19.600	
	47							1.00 30.05
MOTA		CB	PHE	9	27.309	12.376	41.820	1.00 16.95
ATOM	48	CG	PHE	3	26.222	13.355	+2.163	1.00 13.29
MOTA	49	CD1	PHE	3	25.672	13.378	43.447	1.00 17.27
MOTA	50	CD2	PHE	3	25.726		41.189	1.00 13.12
ATOM	51	CEI		8	24.661	14.290		
ATOM	52					14.250	43.772	
			PHE	3	24.712	-1.127	-1.499	1.00 13.19
MOTA	53	CZ	PHE	3	24.192	15.137	42.794	
ATOM	54	11	THR	3	27.798	9.074	+0.599	1.00 27.35
ATOM	55	Cλ	THR	ä	28.704	8.122	40.175	1.00 34.93
ATOM	56	C	THR	. 9	28.709	-,999	12.636	1.00 45.11
ATOM	57	0		é	20.709			1.00 45.11
			THR	G	39.642	7.452	18.062	1.00 50.55
ATOM	58	C3	THR	9	28.447	6.795	40.892	44.00
ATOM	50	OG 1	THR	9	19.619	1,330	+1.527	1.00 40.40
ATOM	50	CGS	THR	;	17.801	1.333	12.250	00 79.75

ATOM	61	N	GLY	10	27.690	8.510	37.956	1.00 30.53
ATOM	6.2	CA	GLY	10	27.689	8.458	36.507	1.00 23.21
ATOM	63	c	GLY	10	27.144	9.746	35.914	1.00 16.55
ATOM	64	ŏ	GLY	10	27.011	10.729	36.617	1.00 25.70
ATOM	65	и	VAL	11	26.835	9.719	34.629	1.00 16.39
ATOM	66	CA	VAL	11	26.209	10.863	33.971	1.00 22.28
ATOM	67	С	VAL	11	24.758	11.020	34.479	1.00 29.60
ATOM	68	0	VAL	11	23.972	10.062	34.456	1.00 20.43
ATOM	69	CB	VAL	11	26.173	10.664	32.467	1.00 30.87
ATOM	70	CG1	VAL	11	25.912	11.980	31.734	1.00 31.75 1.00 33.85 1.00 20.12
ATOM	71	CG 2	VAL	11	27.480	10.048	32.015	1.00 33.85
ATOM	72	N	VAL	12	24.417	12.227	34.931	1.00 20.12
ATOM	73	CA	VAL	12	23.080	12.561	35.433	1.00 12.88
	74	C		12				
ATOM	75	0	VAL		22.407	13.624	34.516	
ATOM			VAL	12	23.007	14.639	34.179	1.00 13.42
ATOM	76	CB	VAL	12	23.270	13.077	36.839	1.00 15.01
ATOM	77	CG1	VAL	12	22.000	13.662	37.422	1.00 17.57 1.00 16.55 1.00 14.72
ATOM	78	CG2	VAL	12	23.781	11.936	37.728	1.00 16.55
ATOM	79	N	PRO	13	21.180	13.382	34.066	1.00 14.72
ATOM	80	CA	PRO	13	20.493	14.382	33.265	.00 10.76
ATOM	81	C	PRO	13	20.116	15.589	34.141	1.00 7.65
ATOM	82	õ	PRO	13	19.797			. 00 15 14
						15.468	35.337	1.00 15.14
MOTA	83	CB	PRO	13	19.225	13.707	32.745	
ATOM	84	CG	PRO	13	19.043	12.422	33.550	1.00 19.69
ATOM	85	CD	PRO	13	20.315	12.195	34.340	1.00 15.41
ATOM	86	11	ILE	14	20.196	16.766	33.557	1.00 19.69 1.00 15.41 1.00 14.91
MOTA	87	CA	ILE	14	19.893	17.991	34.266	1.00 12.93
ATOM	88	C	ILE	14	18.768	13.760	33.596	1.00 12.08
MOTA	89	0	ILE	14	18.724	18.878	32.399	1.00 11.04
ATOM	90	CB	ILE	14	21.109	18.905	34.325	1.00 12.93 1.00 12.08 1.00 11.04 1.00 16.54 1.00 18.08 1.00 11.56
ATOM	91	CG1	ILE	14	22.271	18.169	35.015	1.00 18.08
ATOM	92	CG2	ILE	14	20.783	20.207		1.00 11.56
							35.084	00 11.56
MOTA	93	CD1	ILE	14	23.642	18.836	34.738	1.00 10.13
MOTA	94	И	LEU	15	17.899	19.307	34.421	1.00 13.85
ATOM	95	CA	LEU	15	16.811	20.136	33.955	1.00 14.82 1.00 3.62 1.00 10.00
MOTA	96	C	LEU	15	16.915	21.474	34.685	1.00 3.62
ATOM	97	0	LEU	15	17.080	21.509	35.901	1.00 10.00
ATOM	99	CB	LEU	15	15.462	19.450	34.285	1.00 21.25
ATOM	99	CG	LEU	15	14.412	19.541	33.199	
ATOM	100	CD1	LEU	15	13.279	20.440	33.679	1.00 46.97
ATOM	101	CD2	LEU	15	15.008	20.098	31.913	1.00 49.22
ATOM	102	N	VAL	16		22.556		
					16.885		33.919	1.00 10.56
ATOM	103	CA	VAL	16	16.964	23.905	34.479	1.00 10.23
MOTA	104	С	VAL	16	15.716	24.727	34.063	2.00 9.47
ATOM	105	0	VAL	16	15.347	24.748	32.904	1.00 16.72
MOTA	106	CB	VAL	16	18.273	24.668	34.098	1.00 12.85
ATOM	107	CG1	VAL	16	18.226	26.075	34.691	1.00 12.58
ATOM	108	CG 2	VAL	16	19.520	23.945	34.628	1.00 14.24
ATOM	109	11	GLU	17	15.059	25.317	35.060	1.00 14.43
ATOM	110	CA	GLU	: 7	13.904	26.144	34.870	1.00 14.43
ATOM	111	c	GLU	17	14.086	27.474	35.571	1.00 9.38
ATOM	112	ŏ	GLU	Ξź	14.331	27.524	26.765	1.00 9.38
ATOM	113	CB	GLU	Ξź				
					12.650	25.402	35.344	
ATOM	114	CG	GLU	1.7	12.436	24.178	34.447	1.00 15.37
ATOM	115	CD	GLU	17	11.865	24.573	33.105	1.00 49.50
MOTA	116		, GTD	1.7	11.160	25.557	32.950	1.00 83.46
ATOM	117	OE 2	GLU	17	12.220	23.766	32.127	1.00 38.75
ATOM	118	N	LEU	1.8	13.990	28.571	34.805	1.00 17.82
ATOM	119	CA	LEU		14.116	29.914	35.401	1.00 15.51
ATOM	120	C	LEU	18 18 18	12.962	30.855	35.057	1.00 14.91
ATOM	121	0	LEU		12.585	30.978		
ATOM		CB					33.917	1.00 14.31
	122		LEU	-3	15.426	30.630	35.005	1.00 13.56
ATOM	123	CG	LEU	73	15.533	32.049	35.579	1.00 19.27
ATOM	124	CD1	LEU		16.740	32.182	36.489	1.00 21.40
ATOM	125	CD2	LEU	: 3	15.682	33.033	34.438	1.00 12.38
MOTA	126	11	ASP	19	12.480	31.551	35.082	1.00 17.88
ATOM	127	CA	ASP	19	11.476	32.577	35,940	1.00 19.57

ATOM	:28	С	ASP	19	12.098	33.896	36.360	1.00 11.65
ATOM	129	õ	ASP	19	12.486	34.044	37.493	1.00 16.52
ATOM	130	CB	ASP	19	10.234	32.305	36.847	1.00 24.92
ATOM	131	CG	ASP	19	9.305	31.262	36.282	1.00 38.46
ATOM	132	OD1	ASP	19	8.572	30.587	36.989	1.00 61.49
ATOM	133	OD2	ASP	19	9.337	31.189	34.949	1.00 22.44
ATOM	134	N	GLY	20	12.178	34.863	35.471	1.00 15.32
ATOM	135	CA	GLY	20	12.784	36.101	35.908	1.00 19.52
ATOM	136	С	GLY	20	12.048	37.385	35.538	1.00 19.35
ATOM	137	0	GLY	20	11.240	37.443	34.628	1.00 18.22
ATOM	138	N	ASP	21	12,401	38.407	36.286	1.00 13.19
ATOM	139	CA	ASP	21	11.908	39.737	36.112	1.00 15.36
ATOM	140	С	ASP	21	13.039	40.683	36.424	1.00 12.77
ATOM	141	0	ASP	21	13.517	40.742	37.569	1.00 15.18
ATOM	142	CB	ASP	21	10.701	40.036	37.040	1.00 22.26
ATOM	143	CG	ASP	21	10.230	41.491	37.022	1.00 30.80
MOTA	144	OD1	ASP	21	10.878	42.407	36.557	1.00 27.40
ATOM	145	OD2	ASP	21	9.062	41.658	37.604	1.00 45.92
ATOM	146	N	VAL	22	13.464	41.393	35.397	1.00 19.66
MOTA	147	CA	VAL	22	14.524	42.388	35.542	1.00 25.10
ATOM	148	С	VAL	22	14.010	43.780	35.154	1.00 18.25
MOTA	149	0	VAL	22	13.769	44.062	33.955	1.00 15.10
MOTA	150	CB	VAL	22	15.803	42.012	34.750	1.00 26.37
ATOM	151	CG1	VAL	22	16.861	43.127	34.896	1.00 24.27
ATOM	152	CG2	VAL	22	16.365	40.710	35.297	1.00 22.98
MOTA	153	N	ASN	23	13.823	44.641	36.166	1.00 25.32
MOTA	154	CA	ASN	23	13.319	45.993	35.908	1.00 32.81
ATOM	155	C	ASN	23	11.987	45.958	35.142	00 32.77
MOTA	156	0	ASN	23	11.774	46.730	34.187	1.00 30.47
MOTA	157	CB	ASN	23	14,344	46.831	35.096	1.00 31.26
ATOM ATOM	158 159	CG OD1	ASN	23	15.374	47.607	35.938	1.00 24.72
ATOM	160	ND2	ASN	23	15.795	47.183 48.723	37.024	1.00 27.22
ATOM	161	ND2	ASN	24	15.829	45.024	35.389 35.519	1.00 41.15
ATOM	162	CA	GLY	24	9.831	44.919	34.848	1.00 24.95
ATOM	163	c	GLY	24	9.832	44.111	33.573	1.00 23.22
ATOM	164	ŏ	GLY	24	8.780	43.868	33.024	1.00 28.37
ATOM	165	14	HIS	25	11.000	43.691	33.071	1.00 28.37
ATOM	166	CA	HIS	25	11.042	42.840	31.877	1.00 19.30
ATOM	167	C	HIS	25	10.981	41.373	32.316	1.00 19.30 1.00 27.26 1.00 25.47
MOTA	168	0	HIS	25	11.898	40.850	32.951	1.00 25.47
ATOM	169	CB	HIS	25	12,268	43.060	30.958	1.00 24.20
ATOM	170	CG	HIS	25	12.313	44.382	30.218	1.00 24.20 1.00 33.04 1.00 37.58 1.00 42.76 1.00 39.14
ATOM	171	ND1	HIS	2.5	12.917	45.514	30.758	1.00 37.58
ATOM	172	CD2	HIS	25	11.876	44.716	28.971	1.00 42.75
ATOM	173	CEI	HIS	25	12.801	46.497	29.867	1.00 39.14
ATOM	174	NE2	HIS	25	12.185	46.050	28.778	1.00 42.80
ATOM	175	И	LYS	26	9.872	40.728	32.028	1.00 25.90
ATOM	176	CA	LYS	26	9.675	39.355	32.446	1.00 26.27 1.00 27.09 1.00 25.75
ATOM	177	С	LYS	26	10.154	38.361	31.429	1.00 27.09
ATOM	178	0	LYS	26	10.027	38.576	30.232	1.00 25.75
ATOM	179	CB	LYS	26	8.230	39.069	32.863	1.00 27.58
ATOM	180	CG	LYS	26	7.873	39.770	34.166	1.00 44.94
ATOM	181	CD	LYS	26	6.369	39.914	34.400	1.00 71.44
ATOM	182	CE	LYS	26	6.008	41.000	35.421	1.00 45.29 1.00 22.04 1.00 15.78
ATOM	183	N	PHE	27	10.703	37.250	21.910	1.00 22.04
ATOM	184	CA	PHE	27	11.164	36.236	30.978	
ATOM	185	C	PHE	27	11.273	34.863	31.619	1.00 14.75
ATOM	186	O CB	PHE	27	11.293	34.722	32.842	1.00 18.94
ATOM	187 188	CG	SHE	27	12.495	36.638	30.287	1.00 21.58
ATOM	189	CD1	PHE	27 27	13.599	36.826	31.311	1.00 22.06
ATOM ATOM	190	CD2	PHE	27	14.490	35.791	31.612	1.00 15.94 1.00 21.58 1.00 22.06 1.00 23.61
ATOM	190	CEI	PHE	27.	15.487	35.963		1.00 17.55
ATOM	192	CE2	PHE	27	14.747	38.234	32.579 32.931	1.00 14.41
ATOM	193	CZ	PHE	17	15.621	37.187	33.234	1.00 19.75
ATOM	194	N	SER	23	11.370	33.857	33.234	1.00 12.40
						-3.007	.0.752	

ATOM	195	CA	SER	28	11.492	32.479	31.186	1.00 15.59
ATOM	196	С	SER	28	12.579	31.749	30.379	1.00 15.96
ATOM	197	0	SER	28	12.699	31.933	29.167	1.00 18.99
ATOM	198	CB	SER	28	10.143	31.702	31.086	1.00 14.48
ATOM	199	OG	SER	28	9.510	31.678	32.353	1.00 31.95
ATOM	200	21	VAL	29	13.335	30.902	31.073	1.00 16.73
ATOM	201	CA	VAL	29	14.361	30.093	30.435	1.00 14.06
ATOM ATOM	202 203	C	VAL	29 29	14.258	28.614	30.817	1.00 6.80
ATOM	203	CB	VAL	29	15.768	30.570	31.987	1.00 10.85
ATOM	205	CG1	VAL	29	16.826	29.599	30.234	1.00 15.30
ATOM	206	CG2	VAL	29	15.989	32.001	30.357	1.00 16.37
ATOM	207	N	SER	30	14.462	27.781	29.824	1.00 11.31
ATOM	208	CA	SER	30	14.535	26.351	30.011	1.00 17.96
ATOM	209	C	SER	30	15.917	25.818	29.571	1.00 11.26
ATOM	210	0	SER	30	16.398	26.157	28.513	1.00 13.17
ATOM	211	CB	SER	30	13.471	25.603	29.202	1.00 19.91
ATOM	212	OG	SER	30	12.249	25.667	29.882	1.00 48.74
ATOM	213	N	GLY	31 31	16.480	24.926	30.364	1.00 9.88
ATOM ATOM	214	CA	GLY	31	17.718	24.321	29.977	1.00 12.44
ATOM	216	C	GLY	31	17.737 17.149	22.816	30.249	1.00 13.16
ATOM	217	n	GLU	32	18.459	22.112	29.433	1.00 13.44
ATOM	218	CA	GLU	32	18.622	20.670	29.570	1.00 13.73
ATOM	219	С	GLU	32	20.079	20.297	29.262	1.00 17.33
ATOM	220	0	GLU	32	20.734	20.946	28.456	1.00 15.56
ATOM	221	CB	GLU	32	17.761	19.893	28.543	1.00 12.67
ATOM	222	CG	GLU	32	16.264	20.187	28.618	1.00 26.43
MOTA	223	CD	GLU	32	15.501	19.547	27.468	1.00 21.13
ATOM	224	OE1	GLU	32	15.996	18.767	26.698	1.00 23.45
ATOM	225	0E2	GLU	32	14.292	20.022	27.337	1.00 30.63
ATOM	226	CA	GLY	33 33	20.534 21.860	19.207	29.822	1.00 15.36
ATOM ATOM	227 228	CA	GLY	33	22.236	18.687	29.518	1.00 12.84
ATOM	229	ŏ	GLY	33	21.390	16.919	31.011	1.00 13.56
ATOM	230	11	GLU	34	23.525	17.453	30.702	1.00 15.15
ATOM	231	CA	GLU	34	23.971	15.450	31.621	1.00 18.14
ATOM	232	C	GLU	34	25.220	16.874	32.367	1.00 16.26
ATOM	233	0	GLU	34	25.926	17.760	31.944	00 18.6/
ATOM	234	CB	GLU	34	24.180	15.114	30.927	1.00 22.53
ATOM	235	CG	GLU	34	24.948	15.261	29.624	1.00 33.78
ATOM	236	CD	GLU	34	24.879	14.020	28.796	1.00 55.15
ATOM ATOM	237 238	OE1	GLU	34 34	25.861	13.719	28.534	1.00 45.39
ATOM	239	N N	GLY	35	25.461	16.222	28.430 33.485	1.00 11.20
ATOM	240	CA	GLY	35	26.611	16.502	34.315	1.00 10.62
ATOM	241	c	GLY	35	27.293	15.192	34.662	1.00 19.92
ATOM	242	ŏ	GLY	35	26.650	14.161	34.750	1.00 16.69
ATOM	243	21	ASP	36	28.594	15.238	34.860	1.00 16.92
ATOM	244	CA	ASP	36	29.367	14.061	35.221	1.00 16.19
ATOM	245	С	ASP	36	30.396	14.505	36.233	1.00 13.94
ATOM	246	0_	ASP	36	31.469	15.004	35.879	1.00 15.77
ATOM	247	CB	ASP	36	30.032	13.457	33.948	1.00 19.98
ATOM ATOM	248	CG OD 1	ASP	36 36	30.681	12.066	34.075	1.00 31.92
ATOM	250		ASP	36	31.236	11.519	33.141	1.00 30.97
ATOM	251	11	ALA	37	30.015	11.402	37.490	1.00 13.40
ATOM	252	CA	ALA	37	30.818	14.846	38.582	1.00 12.98
ATOM	253	¢	ALA	37	32.181	14.145	38.637	1.00 21.94
ATOM	254	ō	ALA	37	33.084	1.604	39.331	1.00 13.61
ATOM	255	CB	ALA	37	30.070	14.741	39.916	1.00 11.49
ATOM	256	::	THR	38	32.307	13.016	37.945	
ATOM	257	CA	THR	38	33.581	12.290	37.943	1.00 19.94
ATOM	258	С	THR	38	34.705	13.114	37.335	1.00 19.94 1.00 25.61 1.00 17.89
MOTA	259	0	THR	38	35.850	13.069	37.775	1.00 17.29
ATOM	260	CB	THR	38	33.462	10.398	37.299 38.067	1.00 22.57
ATOM	261	OG 1	. T. K	23	32.543	0	25.06/	1.00 29.86

ATOM	262	CG2	THR	38	34.821	10.213	37.355	1.00 22.90
ATOM	263	11		39			36.347	1.00 18.45
			TYR		34.323	13.920		1.00 10.45
MOTA	264	CA	TYR	39	35.210	14.837	35.675	1.00 9.39
MOTA	265	С	TYR	39	34.874	16,291	35.991	1.00 14.41
ATOM	266	0	TYR	39	35.454	17.177	35.410	1.00 16.24
ATOM	267	CB	TYR	39	35.156	14.582	34.180	1.00 16.24
ATOM	268	CG	TYR	39 .	35.426	13.137	33.929	1.00 28.73
ATOM	269	CD1	TYR	39		12.633	34.065	1.00 33.75
					36.715			
ATOM	270	CD2	TYR	39	34.392	12.249	33.642	1.00 39.19
ATOM	271	CEI	TYR	39	36.982	11.276	33.828	1.00 29.75
ATOM	272	CE2	TYR	39	34.635	10.885	33.435	1.00 45.41
ATOM	273	CZ	TYR	39	35.943	10.410	33.570	1.00 57.62
ATOM	274	OH	TYR	39	36.199	9.070	33.364	1.00 70.77
ATOM	275	11	GLY	40	33.935	16.525	36.929	1.00 9.94
ATOM	276	CA	GLY	40	33.474	17.879	37.266	1.00 7.02
ATOM	277	c	GLY	40	32.952	18.600	36.004	1.00 9.45
ATOM	278	o	GLY	40	33.068		35.829	1.00 12.63
						19.830		
ATOM	279	N	LYS	41	32.380	17.823	35.092	1.00 5.44
ATOM	280	CA	LYS	41	31.954	18.335	33.842	1.00 6.63
ATOM	281	C	LYS	41	30.414	18.554	33.703	1.00 20.92
ATOM	282	0	LYS	41	29.617	17.693	34.085	1.00 12.94
ATOM	283	CB	LYS	41	32.360	17.357	32.827	1.00 8.27
ATOM	284	CG	LYS	41	32.099	17.771	31.419	1.00 13.19
ATOM	285	CD	LYS	41	32.521	16.644	30.481	1.00 20.20
ATOM	286	CE	LYS	41	32.690	17.068	29.032	1.00 35.79
ATOM	287	NZ	LYS	41	33.113	17.060	28.147	1.00 47.56
						15.954		
ATOM	288	21	LEU	42	30.049	19.684	33.069	1.00 18.31
ATOM	289	CA	LEU	42	28.643	20.064	32.794	1.00 16.08
ATOM	290	C	LEU	42	28.456	20.422	31.330	1.00 16.08
ATOM	291	0	LEU	42	29.240	21.168	30.787	
ATOM	292	CB	LEU	42	28.223	21.300	33.621	1.00 13.22
ATOM	293	CG	LEU	42	28.007	21.061	35.082	1.00 16.70
ATOM	294	CD1	LEU	42	27.894	22.406	35.782	1.00 13.79
ATOM	295	CD2	LEU	42	26.732	20.243	35.295	1.00 18.70
ATOM	296	N	THR	43	27.395	19.914	30.672	1.00 8.04
								1.00 6.04
ATOM	297	CA	THR	43	27.103	20.275	29.282	1.00 4.87 1.00 17.23 1.00 14.38
MOTA	298	С	THR	43	25.636	20.666	29.186	1.00 17.23
ATOM	299	0	THR	43	24.811	19.818	29.442	1.00 14.38
ATOM	300	CB	THR	43	27.351	19.140	28.317	1.00 21.59
ATOM	301	OG1	THR	43	28.692	18.743	23.415	1.00 42.74
ATOM	302	CG2	THR	43	27.073	19.675	26.917	1.00 31.23
ATOM	303	77	LEU	44	25.327	21.934	13.830	1.00 11.83
ATOM	304	CA	LEU	44	23.944	22.409	13.830 13.847 17.668	1.00 13.81
ATOM	305	c	LEU	44	23.589	23.307	7 668	1.00 18.19
ATOM	306	ŏ	LEU	44	24.416	23.989	27.107	1.00 13.86
ATOM	307	CB	LEU	44	23.725	23.275	30.125	1.00 15.37
ATOM	308	CG	LEU	44	23.369	22.584	31.456	1.00 24.69
ATOM	309	CD1	LEU.	44	21.869	22.381	31.601	1.00 23.20
ATOM	310	CD2	LEU	44	24.083	21.286	31.650	1.00 46.18
ATOM	311	N	LYS	45	22.294	23.331	27.339	1.00 10.29
ATOM	312	CA	LYS	45	21.752	24.224	15.358	1.00 11.94
ATOM	313	c	LYS	45	20.534	24.913	25.957	1.00 11.94
ATOM	314	ō	LYS	45	19.665	24.248		1.00 18.43
ATOM	315	СВ	LYS	45	21.409	23.560	060	1.00 13.75
ATOM	316	CG		45	20.878	24.556	27.530 25.060 24.045	1.00 13.75 1.00 8.83
	317	CD	LYS LYS	45	20.486	24.550	24.045	1.00 8.83
MOTA			LIS			23.863	22.746	1.00 26.87
MOTA	318	CE	LYS	45	19.574	24.688	11.842	1.00 16.58
ATOM	319	210	LYS	45	19.318	24.024	20.555	1.00 26.87 1.00 16.58 1.00 18.33 1.00 12.34 1.00 13.32
MOTA	320	11	PHE	46	20.535	26.236	25.910	1.00 12.34
ATOM	321	CA	PHE	46	19,463	27.048	17.451	
ATOM	322	C	PHE	46	18.759	27.718	16.343	1.00 18.26
ATOM	323	9	PHE	+6	19.386	28.093		
ATOM	324	CB	PHE	46	19.934	33.101	13.473	1.00 15.29
ATOM	325	CG	PHE	÷6\	20.773	27.495	2 662	1.00 13.91
ATOM	326	CD1	PHE	46	22.132		-7.554	1.00 17.06
ATOM	327	502		46		37.268	-2-23/	17.06
					20.209	27.121 26.693	18.473 19.552 19.337 19.774	1.00 16.83 1.00 15.29 1.00 13.81 1.00 17.06 1.00 8.24
MOTA	328	CEI	PHE	<b>÷</b> 6	22.924	16.693	23.331	1.00 15.95

ATOM	329	CE2	PHE	46	20.979	26.524	31.767	1.00 11.90
ATOM	330	CZ	PHE	46	22.340	26.309	31.540	
ATOM	331	N	ILE	47	17.440	27.845	26.498	1.00 13.24
ATOM	332	CA	ILE	47	16.588	28.453	25.479	1.00 13.24
ATOM	333	C	ILE	47	15.645	29.460	26.118	1.00 20.14
ATOM	334	0	ILE	47	15.039	29.162	27.148	1.00 17.67
ATOM	335	CB	ILE	47	15.737	27.386	24.801	1.00 17.67 1.00 22.67 1.00 20.66
ATOM	336	CG1	ILE	47	16.585	26.271	24.291	1.00 20.66
ATOM	337	CG2	ILE	47	15.024	28.002	23.641	1.00 33.79
ATOM	338	CD1	ILE	47	16.639	26.293	22.805	1.00 23.69
ATOM	339	N	CYS	48	15.564	30.653	25.561	1.00 14.68
MOTA	340	CA	CYS	48	14.681	31.635	26.170	1.00 16.93
ATOM	341	C	CYS	48	13.323	31.352	25.628	1.00 24.18
ATOM	342	0	CYS	48	13.122	31.513	24.453	1.00 20.63
ATOM	343	CB	CYS	48	15.063	33.116	25.885	1.00 16.85
ATOM	344	SG	CY5	48	13.913	34.268	26.712	1.00 22.06
ATOM	345	N	THR	49	12.424	30.871	26.484	1.00 27.31
ATOM	346	CA	THR	49	11.101	30.458	26.042	1.00 32.18
ATOM	347	c	THR	49	10.106	31.572	25.803	1.00.37.51
ATOM	348	O CB	THR	49 49	9.150	31.407	25.061	1.00 35.71
ATOM ATOM	349 350	OG1	THR	49	10.537	29.417	26.972	1.00 23.66
ATOM ·	351	CG2	THR	49	10.387	29.989	28.258 27.022	1.00 30.10
ATOM	352	N N	THR	50	10.314	32.693	26.447	
ATOM	353	CA	THR	50	9.416	33.810	26.283	1.00 32.34
ATOM	354	c	THR	50	9.836	34.711	25.125	1.00 37.98
ATOM	355	ò	THR	50	9.228	35.763	24.904	1.00 37.56
ATOM	356	СЗ	THR	50	9.251	34.611	27.589	1.00 36.23
ATOM	357	OG1	THR	50	10.512	34.980	28.118	1.00 35.37
ATOM	358	CG2	THR	50	8.507	33.773	28.602	1.00 35.37
ATOM	359	N	GLY	51	10.881	34.282	24.372	1.00 31.04
ATOM	360	CA	GLY	51	11,394	35.059	23.239	1.00 32.42
ATOM	361	С	GLY	51	12.865	35.542	23.427	1.00 48.45
ATOM	362	0	GLY	51	13.779	34.737	23.701	1.00 57.11
MOTA	363	N	LYS	52	13.087	36.862	23.282	1.00 35.08
ATOM	364	CA	LYS	52	14.416	37.460	23.416	1.00 35.75
ATOM	365	C	LYS	52	14.827	37.726	24.851	1.00 29.65
ATOM	366	0	LYS	5 2	14.140	38.420	25.620	1.00 25.70
ATOM	367	CB	LYS	5.2	14.577	38.714	22.582	1.00 43.37
ATOM	368	CG	LYS	52	15.772	38.649	21.644	1.00 78.17
ATOM	369	N	LEU	5.3	15.983	37.190	25.250	1.00 19.22
ATOM	370	CA	LEU	53	16.439	37.430	26.596	1.00 13.52
ATOM	371 372	C	LEU	53 53	16.717	38.932	26.775	1.00 17.76
ATOM	373	CB	LEU	53	17.392	39.539	25.973	1.00 21.59
ATOM ATOM	374	CG	LEU	53	17.705 18.100	36.567	26.845	1.00 17.39
ATOM	375	CD1	LEU	53	17.048	36.435 35.621	28.302 29.053	1.00 17.43
MOTA	376	CD2	LEU	53	19.440	35.718	28.368	1.00 20.12
ATOM	377	N	PRO	54	16.197	39.525	27.817	1.00 16.69
ATOM	378	CA	PRO	54	16.324	40.962	28.092	1.00 18.60
ATOM	379	c	PRO	54	17.638	41.414	28,707	1.00 25.39
ATOM	380	ŏ	PRO	54	17.865	42.609	28.861	1.00 18.88
ATOM	381	CB	PRO	54	15.268	41.265	29.139	1.00 22.52
ATOM	382	CG	PRO	54	14.832	39.933	29.720	1.00 26.02
ATOM	383	CD	PRO	54	15.318	38.855	28,779	1.00 21.26
ATOM	384	71	'VAL	5.5	18.435	40.455	29.161	1.00 23.32
ATOM	385	CA	VAL	5.5	19.746	40.716	29.711	1.00 15.83
MOTA	386	C	VAL	55	20.688	39.868	28.973	1.00 19.38
ATOM	387	0	VAL	55	20.268	39.035	28.219	1.00 20.34
ATOM	388	CB	VAL	55	19.814	40.409	31.147	1.00 17.67
ATOM	389	CG1	VAL	55	18.864	41.340	31.851	1.00 22.52
MOTA	390	CG2	VAL	55	19.402	38.959	31.397	2.00 17.11
MOTA	391	11	PRO	56	21.963	40.070	29.167	
ATOM	392	CA	PRO	- 5 - 5	22.911	39.258	28.447	1.00 13.09
ATOM	393	C	PRO	5.6	23.059	37.834	29.038	1.30 5.83
ATOM ATOM	394 295	O CB	PRO	5.5 5.5	23.067	37.631	30.254	1.00 12.35
" TON	232	CB	PRO	- 0	24.231	40.062	23.420	1.00 13.34

110 3								
ATOM	396	CG	PRO	56	23.851	41.478	28.849	1.00.20.73
ATOM	397	CD	PRO	5.5	22.525	41.379	29.578	1.00 20.73
ATOM	398	U	TRP	57	23.202	36.848	28.158	1.00 11.12
ATOM	399	CA	TRP	57	23.354	35.458	28.595	1.00 12.55
ATOM	400	C	TRP	57	24.411	35.239	29.700	1.00 14.13
ATOM	401	0	TRP	57	24.178	34.586	30.709	1.00 11.49
MOTA	402	CB	TRP	57	23,604	34.535	27.406	1.00 10.56
MOTA	403	CG	TRP	57	22.335	34.237	26.641	1.00 12.55
ATOM	404	CD1	TRP	57	21.999	34.714	25.426	1.00 16.24
ATOM	405	CD2	TRP	57	21.281	33.327	27.013	1.00 12.50
ATOM	406	NE1	TRP	57	20.784	34.200	25.018	1.00 14.25
ATOM	407	CE2	TRP	57	20.315	33.354	25.963	1.00 14.65
ATOM	408 409	CE3	TRP	57	21.052	32.521	28.129	1.00 12.01
ATOM ATOM	410	CZ2	TRP	57 57	19.148 19.887	32.583	26.007	1.00 14.36
ATOM	411	CH2	TRP	57	18.945	31.767 31.818	28.170 27.128	1.00 14.23
ATOM	412	N	PRO	58	25.594	35.800	29.518	1.00 15.78
ATOM	413	CA	PRO	58	26.629	35.616	30.503	1.00 9.53
ATOM	414	č	PRO	58	26.241	36.010	31.878	1.00 9.71
ATOM	415	ō	PRO	58	26.760	35.467	32.825	1.00 11.70
ATOM	416	CB	PRO	58	27.833	36.441	30.040	1.00 11.70
ATOM	417	CG	PRO	58	27.597	36.748	28.582	1.00 18.50
MOTA	418	CD	PRO	58	26.137	36.432	28.278	1.00 15.82
ATOM	419	N	THR	59	25.336	36.977	32.021	1.00 7.54
ATOM	420	CA	THR	59	24.976	37.366	33.357	1.00 4.53
ATOM	421	c	THR	59	24.228	36.258	34.137	1.00 8.41
	422	0	THR	59	24.174	35.251	35.367	1.00 10.57
ATOM ATOM	423 424	CB OG1	THR	59 59	24.197	28.691 38.480	33.384 32.844	1.00 15.54
ATOM	425	CG2	THR	59	22.895	39.731	32.542	1.00 15.76
ATOM	426	N	LEU	60	23.686	25.304	33.427	1.00 11.99
ATOM	427	ĊA	LEU	60	22.899	34.248	34-073	1.00 9.15
ATOM	428	С	LEU	60	23.657	32.944	34.385	1.00 15.62
ATOM	429	0	LEU	60	23.118	32.027	35.042	1.00 11.99
A <b>TO</b> M	430	CB	LEU	60	21.645	33.914	33.203	1.00 7.67
ATOM	431	CG	LEU	60	20.728	35.111	33.042	1.00 14.05
ATOM	432	CD1	LEU	60	19.620	34.775	32.062	1.00 14.54
ATOM	433	CD2	LEU	60	20.142	35.456	34.394	1.00 10.67
ATOM ATOM	434 435	N CA	VAL	61 61	24.893	22.837	33.917	1.00 11.27
ATOM	436	C	VAL	61	25.656 25.678	31.587	34.094	1.00 4.37 1.00 6.02 1.00 10.78 1.00 7.14
ATOM	437	ŏ	VAL	61	25.355	29.805	35.743	1.00 10.75
ATOM	438	CB	VAL	61	27.050	31.643	33.406	1.00 7.14
ATOM	439	CG1	VAL	61	27.888	30.396	33.805	1.00 6.47
ATOM	440	CG2	VAL	51	26.890	31.745	31.876	1.00 6.63
ATOM	441	N	THR	52	26.053	31.843	36.442	1.00 7.02
ATOM	442	CA	THR	62	26.178	31.421	37.808	1.00 6.51
MOTA	443	С	THR	62	24.862	30.954	38.410	1.00 9.22
ATOM	444	O CB	THR	62	24.801	30.163	39.352	1.00 6.99
ATOM ATOM	445	OG1	THR	62 62	26.816	32.520 33. <b>744</b>	38.660	1.00 16.97
ATOM	447	CG2	THR	62	26.103 28.297	32.708	38.453	1.00 12.00 1.00 S.86
ATOM	448	N	THR	63	23.814	31.547	37.910	1.00 9.98
ATOM	449	CA	THR	63	22.457	31.212	38.388	1.00 6.69
ATOM	450	C	THR	63	22.033	29.830	37.865	1.00 8.14
ATOM	451	0	THR	63	21.499	23.984	38.604	1.00 13.48
ATOM	452	CB	THR	53	21.458	32.312	37.925	1.00 11.14
ATOM	453	OG1	THR	63	21.785	33.498	38.602	1.00 11.75
ATOM	454	CG2	THR	53	20.024	31.897	38.296	1.00 9.31
ATOM	455	N	PHE	54	22.250	29.620	36.583	1.00 10.19
ATOM ATOM	456 457	CA .	PHE	64 64	21.895	28.371	15.995	1.00 8.00
ATOM	458	0	PHE	54	22.774	27.253	36.518	1.00 25.25
ATOM	459	CB	PHE	54 54	22.313	28.147	36.761 34.513	1.00 9.54
ATOM	460	CG	PHE	64	21.233	29.357	33.750	1.00 10.96
ATOM	461	CD1	PHE	54	21.724	19.954	12.593	1.00 7.15
ATOM	462	CD2	PHE	54	19.899	19.563	34.106	1.00 6.88 1.00 10.96 1.00 9.15 1.00 14.43

ATOM	463	CEI	PHE	64	20.936	30.792	31.805	1.00 14.20
ATOM	464	CE2	PHE	64	19.077	30.375	33.317	1.00 13.95
ATOM	465	CZ	PHE	64	19.597	30.983	32.171	1.00 16.35
HETATM	466	21	CRO	66	24.077	27.513	36.610	1.00 11.86
HETATM	467	CG1	CRO	66	25.155	25.422	34.796	1.00 16.67
HETATM	468	OG1	CRO	66	26.679	27.129	35.461	1.00 14.22
HETATM	469	CBl	CRO	66	25.931	26.035	35.930	1.00 10.77
HETATM	470	CAl	CRO	66	25.011	26.478	37.078	1.00 7.34
HETATM	471	Cl	CRO	66	25.718	26.991	38.253	1.00 17.70
HETATM	472	N2	CRO	66	26.975	27.732	38.216	1.00 9.21
HETATM	473	OH	CRO	66	32.894	30.804	36.971	1.00 13.84
HETATM	474	CD2	CRO	66	30.487	30.110	39.805	1.00 10.79
HETATM	475	CE2	CRO	6 <b>6</b>	31.614	30.563	39.085	1.00 10.01
HETATM	476	CZ	CRO	66	31.718	30.300	37.721	1.00 9.48
HETATM	477	CE1	CRO	66	30.707	29.546	37.033	1.00 17.44
HETATM	478	CD1	CRO	66	29.541	29.103	37.742	1.00 11.31
HETATM	479	CG2	CRO	66	29.437	29.370	39.124	1.00 7.67
HETATM	480	CB2	CRO	66	28.329	28.822	39.960	1.00 10.75
HETATM	481	CA2	CRO	66	27.197	28.245	39.512	1.00 16.08
HETATM HETATM	482 483	C2 O2	CRO	6 <b>6</b> 66	26.043	27.875	40.370	1.00 5.46
HETATM	484	113	CRO	66	26.022 25.240	26.978	39.517	1.00 13.20
HETATM	485	CA3	CRO	66	23.840	26.511	39.734	1.00 10.40
HETATH	486	CA3	CRO	66	23.413	25.550	40.817	1.00 10.40
HETATM	487	03	CRO	66	22.747	26.014	41.764	1.00100.00
ATOM	488	11	VAL	68	23.737	24.208	41.764	1.00 29.95
ATOM	489	ĊA	VAL	68	24.209	22.972	40.304	1.00 17.16
ATOM	490	c	VAL	68	25.692	22.550	40.734	1.00 14.88
ATOM	491	ō	VAL	68	26.378	21.821	40.026	1.00 9.03
ATOM	492	CB	VAL	68	23.870	22.899	38.831	1.00 18.94
ATOM	493	CG1	VAL	68	24.685	22.088	37.942	1.00 17.17
ATOM	494	CG2	VAL	68	22.396	22.538	38.680	1.00 18.80
MOTA	495	22	GLN	69	26.129	22.965	41.914	1.00 11.04
ATOM	496	CA	GLN	69	27.465	22.764	42.394	1.00 15.00
ATOM	497	C	GLN	69	27.749	21.366	42.893	1.00 22.46
ATOM	498	0	GLN	69	28.876	21.025	43.154	1.00 15.84
ATOM	499	CB	GLN	69	27.929	23.852	43.414	1.00 10.93
ATOM	500	CG	GLN	69	29.202	25.174	42.615	1.00 14.13
MOTA	501	CD	GLN	69	28.216	25.385	43.520	1.00 17.01
ATOM	502	OE1	GLN	69	27.433	26.476	44.448	1.00 18.94
ATOM	503	TE2	GLN	69	29.151	27.300	43.241	1.00 8.52
ATOM	504	14	CYS	70	26.703	20.540	42.906	1.00 12.10
ATOM	505 506	CA	CYS	70 70	26.862	19.171	43.287	1.00 11.84
ATOM ATOM	507	CO	CYS	70	27.611	18.391 17.242	42.175	1.00 10.54
ATOM	508	Č8	CYS	70	25.476	18.584	42.367	1.00 14.70
ATOM	509	SG	CYS	70	24.325	19.012	42.251	1.00 15.61
ATOM	510	N	PHE	71	27.801	19.029	41.005	1.00 8.64
ATOM	511	CA	PHE	71	28.525	18.419	39.883	1.00 6.59
ATOM	512	c	PHE	71	30,041	18.754	39.876	1.00 16.43
ATOM	513	0	PHE	71	30.753	13.481	38.916	1.00 13.05
ATOM	514	CB	PHE	71	27.951	18.771	38.523	1.00 7.61
ATOM	515	CG	PHE	71	26.669	13.016	38.303	1.00 14.73
ATOM	516	CD1	PHE	71	26.693	15.642	38.050	1.00 10.34
ATOM	517	CD2		71	25.434	18.660	38.453	1.00 17.14
MOTA	518	CE1		71	25.506	18.931	37.866	1.00 15.09
MOTA	519	CE2		71	24.238	15.598	38.300	1.00 15.09
MOTA	520	CZ	PHE	71	24.282	15.598	37.990	1.00 18.49
ATOM	521	31	SER	72	30.500	19.370	40.938	1.00 13.13
ATOM	522	CA	SER	72	31.889	19.715	41.075	1.00 11.65
ATOM	523	C	SER	72	32.689	13.446	41.357	1.00 14.56
MOTA	524	0	SER	72	32.256	17.566	42.122	1.00 10.90
MOTA	525	CB	SER	7.2	32.075	20.672	42.257	1.00 2.65
MOTA	526 527	OG '}	SER	72	31.361		42.038	1.00 19.29
ATOM ATOM	528	CA	ARG	7.3	33.905	13.358	40.794	1.00 16.27
ATOM	526	C	ARG	- 3	34.695	11.212	41.117	1.00 13.55

ATOM	530	0	ARG	73	36.182	18.376	42.599	1.00 16.14
ATOM	531	СВ	ARG	73	35.694	16.817	40.013	1.00 16.80
ATOM	532	CG	ARG	73	36.549	15.616	40.460	1.00 20.13
ATOM	533	CD	ARG	73	37.489	15.093	39.381	1.00 28.47
ATOM	534	NE	ARG	73	38.743	15.859	39.260	
				73		15.859		
ATOM	535	CZ	ARG		39.756	15.777	40.127	1.00 28.04
ATOM	536	NH1		73	39.688	15.004	41.195	1.00 28.76
ATOM	537		ARG	73	40.865	16.504	39.918	1.00 39.65
ATOM	538	N	TYR	74	35.151	16.561	43.424	1.00 12.05
ATOM	539	CA	TYR	74	35.861	16.659	44.690	1.00 11.57
ATOM	540	С	TYR	74	36.946	15.566	44.721	1.00 25.02
ATOM	541	0	TYR	74	36.658	14.387	44.558	1.00 19.71
ATOM	542	CB	TYR	74	34.978	16.528	45.934	1.00 15.51
ATOM	543	CG	TYR	74	34.395	17.850	46.402	1.00 16.59
ATOM	544	CD1	TYR	74	33.455	18.546	45.631	1.00 14.44
ATOM	545	CD2	TYR	74	34.799	18.399	47.618	1.00 15.94
ATOM	546		TYR	74	32.901	19.756	46.059	1.00 7.99
ATOM	547		TYR	74	34.261	19.612	48.058	1.00 18.29
ATOM	548	cz	TYR	74	33.294	20.276	47.298	1.00 13.87
ATOM	549	OH	TYR	74	32.829	21.507	47.738	1.00 18.39
ATOM	550	N	PRO	75	38.181	15.947	44.902	1.00 19.20
ATOM	551	CA	PRO	75	39.213	14.940	44.995	1.00 18.42
ATOM	552		PRO	75	38.958	13.993	46.175	1.00 18.42
		C						
MOTA	553	0	PRO	75	38.373	14.361	47.174	1.00 11.99
ATOM	554	CB	PRO	75	40.514	15.681	÷5.195	1.00 18.31
MOTA	555	CG	PRO	75	40.242	17.158	44.963	00 24.81
ATOM	556	CD	220	75	38.742	17.306	44.694	1.00 15.41
ATOM	557	N	ASP	76	39.433	12.756	46.038	1.00 18.63
ATOM	558	CA	ASP	76	39.269	11.770	47.062	1.00 16.19
ATOM	559	C	ASP	76	39.581	12.280	48.431	1.00 15.92
MOTA	560	0	ASP	76	38.862	12.042	49.389	1.00 17.35 1.00 18.69
ATOM	561	CB	ASP	76	40.083	10.507	46.790	1.00 18.69
ATOM	562	CG	ASP	76	39.826	9.432	47.825	1.00 24.04
ATOM	563	001	ASP	76	40.523	9.268	48.817	1.00 29.72
ATOM	564		ASP	76	38.732	8.743	47.584	
ATOM	565	N	HIS	77	40.647	12.984	48.561	1.00 40.96 1.00 18.79
ATCM	566	CA	HIS	77	40.978	13.418	49.877	1.00 19.36
ATOM	567	C	HIS	77	40.117	14.507	30.397	1.00 24.57
ATOM	568	õ	HIS	77	40.205	14.826	51.551	1.00 24.37
ATOM	569	CB	HIS	77	42.435	13.806	50.042	1.00 19.84
ATOM	570	CG	HIS	77	42.433	15.035	49.322	
				77				1.00 17.31
ATOM	571		HIS		42.925	15.028	47.953	1.00 21.86
ATOM	572		HIS	77	42.925	15.295	49.774	1.00 18.70
MOTA	573	CEl	HIS	77	43.203	16.289	47.593	1.00 17.49
ATOM	574	NE2	HIS	77	43.213	17.069	48.668	1.00 18.11
ATOM	575	N	MSE	78	39.277	15.069	49.565	1.00 25.36
ATOM	576	CA	MSE	78	38.412	16.140	50.026	1.00 24.65
ATOM	577	C	MSE	78	36.920	15.774	50.066	1.00 26.47
ATOM	578	0	MSE	78	36.070	16.636	50.260	1.00 28.16
MOTA	579	CB	MSE	78	38.596	17.331	49.121	1.00 26.38
ATOM	580	CG	MSE	78	39.803	18.177	49.406	1.00 27.01
ATOM	581	SE	MSE	78	39.987	19.608	48.117	1.00 43.09
MOTA	582	CE	MSE	78	38.874	20.873	49.044	1.00 27.11
ATOM	583	11	LYS	79	36,606	14.509	49.856	1.00 27.11
ATOM	584	CA	LYS	79	35.216	14.061	÷9.853	1.00 21.54
ATOM	585	C	LYS	79	34.406	14.449	51.082	1.00 21.54
ATOM	586	õ	LYS	79	33.186	14.652	11.025	1.00 21.08
ATOM	587	CB	LYS	79	35.152	12.581	49.612	1.00 23.48
ATOM	588	CG	LYS	79	35.859	12.225	48.317	1.00 23.48
ATOM	589	co	LYS	79	35.159	11.134	47.535	2.00 41.09
ATOM	590	CE	LYS	79	35.796	10.881	46.181	2.00 34.66
ATOM	591	11Z	175	79	35.084			53.46
				30		11.549	45.080	1.00 49.53
ATOM	592	::	ARG		35.069	14.542	52.213	1.00 19.77
ATOM	593	CA	ARG	3,0	34.365	874	53.434	1.00 20.13
ATOM	594	C	ARG	30	33.898	16.311	53,481	1.00 20.13
ATOM	595	0	ARG	30	33.251	5.715	34.467	1.00 23.51
ATOM	596	C3	ARG	30	35.155	14.549	14.700	1.00 24.53

ATOM	597	CG	ARG	30	36.204	15.620	55.034	1.00 29.71
ATOM	598	CD	ARG	30	36.964	15.344	56.335	. 00 61 30
ATOM	599	NE	ARG	30	36.551	16.230	57.415	: 00 71 14
ATOM	600	CZ	ARG	50	37.398	16.882	58.192	1.00 71.14
ATOM	601	NH1	ARG	80	38.714	16.758	58.040	1.00100.00
ATOM	602	NH2	ARG	20	36.917	17.679	59.155	1.00 99.06
ATOM	603	N	HIS	21	34.275	17.121	52.473	2.00 18.77
ATOM	604	CA	HIS	91	33.903	18.547	52,499	1.00 19.60
ATOM	605	С	HIS	31	32.841	18.883	51.486	1.00 18.62
ATOM	606	0	HIS	81	32.557	20.043	51.295	: 00 17 76
ATOM	607	CB	HIS	81	35.129	19.472	52,283	1.00 20.39
MOTA	608	CG	HIS	81	36.221	19,224	53.305	1.00 28.02
MOTA	609	ND1	HIS	81	-36.127	19.701	54.618	1.00 30.59
MOTA	610	CD2	HIS	a1	37.392	18.535	53.202	00 29.02 00 26.24
ATOM	611	CEL	HIS	81	37.218	19.308	55.265	1.00 26.24
ATOM	612	NE2	HIS	81	37.991	18.603	54.452	1.00 28.18
MOTA	613	N	ASP	82	32.298	17.843	50.841	1.00 12.20
ATOM	614	CA	ASP	82	31.358	18.011	49.769	1.00 13.24
ATOM	615	C	ASP	82	29.922	18.148	50.259	1.00 24.30
MOTA	616	0	ASP	82	29.175	17.195	50.243	1.00 16.55
MOTA	617	CB	ASP	9.2	31.480	16.917	48.730	1.00 12.23
MOTA	618	CG	ASP	82	30.642	17.209	47.518	1.00 9.92
ATOM	619	OD1	ASP	82	29.870	18.134	47.459	00 20.31
ATOM	620	OD2	ASP	32	30.938	16.466	46.507	1.00 11.12
MOTA	621	N	PHE	83	29.566	19.353	50.705	1.00 23.66
ATOM	622	CA	PHE	83	28.220	19.634	51.201	1.00 20.23
MOTA	623	C	PHE	83	27.154	19.333	50.168	1.00 20.93
ATOM	624	0	PHE	83	26.116	18.733	50.503	1.00 15.97
MOTA	625	CB	PHE	83	28.077	21.106	51.666	1.00 19.59
ATOM	626	CG	PHE	83	26.624	21.613	51.805	1.00 16.91
ATOM	627	CD1		93	25.946	21.498	53.021	1.00 17.76
MOTA	628	CD2	PHE	33	25.968	22.236	50.734	1.00 18.88
MOTA	629 630	CE1	PHE	83	24.635	21.960	53.156	1.00 24.13
MOTA	631	CE2	PHE	83	24.650	22.690	50.840	1.00 19.24
MOTA MOTA	632	N.	PHE	83 64	24.001	22.575	52.068 48.921	1.00 20.67 1.00 14.06 1.00 12.96
ATOM	633	CA	PHE	24	26.515	19.693	47.809	1.00 12.96
ATOM	634	c	PHE	34	25.893	18.332	47.602	1.00 24.96
MOTA	635	Ö	PHE	84	24.674	18.200	47.534	00 21 55
ATOM	636	CB	PHE	84	27.085	20.265	46.513	1.00 24.96 1.00 21.55 1.00 13.44
ATOM	637	CG	PHE	34	27.630	21.645	46.721	1.00 14.27 1.00 15.17 1.00 13.48
ATOM	638	CD1		84	29.001	21.845	46.890	1.00 15.17
MOTA	639	CD2		94	26.781	22.753	46.752	1.00 13.48
ATOM	640	CE1		34	29.520	23.129	47.073	1.00 14.63
ATOM	641	CE2		84	27.276	24.041	46.969	1.00 16.34
ATOM	642	CZ	PHE	34	28.650	24.221	47.137	1.00 15.77
ATOM	643	N	LYS	95	26.738	17.330	47.482	1.00 14.07
MOTA	644	CA	LYS	85	26.294	15.985	47.283	1.00 13.30
MOTA	645	С	LYS	35	25.657	15.371	48.547	1.00 13.43
MOTA	646	0	LYS	35	24.773	14.509	48.429	1.00 18.46
MOTA	647	CB	LYS	35	27.434	15.089	46.757	1.00 17.38
MOTA	648	CG	LYS	35	27.873	15.372	45.323	1.00 13.93
ATOM	649	CD	LYS	35	28.969	14.381	44.888	1.00 13.23
MOTA	650	CE	LYS	85	29.766	14.819	43.662	1.00 10.36
MOTA	651	NZ	LYS	35	30.319	16.185	43.773	1.00 12.92
MOTA	652	21	SER	36	26.119	15.795	49.752	1.00 11.03
MOTA	653	CA	SER	36	25.610	15.267	50.998	1.00 12.09
MOTA	654	C	SER	36	24.156	15.639	51.240	1.00 21.58
MOTA	655	0	SER	86 ⋅	23.452	14.979	52.013	1.00 19.89
ATOM	656	CB	SER	36	26.448	15.661	52.208	1.00 16.45
ATOM	657	OG	SER	3.6	26.308	17.042	52.495	1.00 22.05
MOTA	658	31	ALA	37	23.705	16.698	50.582	00 15.09
ATOM	659	CA	ALA	£7	22.333	17.138	50.762	1.00 19.52
MOTA	560	С	ALA	3.7	21.337	16.399	19.870	1.00 18.60
ATOM	661	0	ALA	3.7	20.162	16.557	30.040	1.00 19.55
ATOM	562	CB	ALA	3.7	22.204	19.647	50.632	1.00 19.23
MOTA	563	::	MSE	5.8	21.835	15.536	48.976	1.00 14.05

MOTA	- 554	CA	MSE	88	21.007	14.796	48.035	1.00 15.32
ATOM	665	C	MSE	88	20.496	13.448	48.579	1.00 21.48
	566					12.876	49.457	1.00 23.03
ATOM		0_	MSE	88	21.109			
ATOM	667	CB	MSE	88	21.848	14.593	46.791	1.00 16.93
ATOM	668	CG	MSE	88	22.263	15.891	46.131	1.00 10.56
ATOM	669	SE	MSE	88	20.737	16.894	45.394	1.00 31.99
MOTA	670	CE	MSE	88	21,318	18.684	45.748	1.00 28.86
ATOM	671	N	PRO	89	19.363	12.930	48.084	1.00 14.78
ATOM	672	CA	PRO	89	18.552	13.475	47.008	1.00 14.80
ATOM	673	C		89	17.572	14.611	47.385	1.00 12.10
			PRO					
ATOM	674	0	PRO	89	17.085	15.301	46.493	1.00 18.06
ATOM	675	CB	PRO	89	17.733	12.294	46.494	1.00 17.00
MOTA	676	CG	PRO	89	17.726	11.261	47-607	1.00 15.23
ATOM	677	CD	PRO	89	18.844	11.642	48.560	1.00 17.16
MOTA	678	N	GLU	90	17.278	14.795	48.695	1.00 14.63
ATOM	679	CA	GLU	90	16.348	15.838	49.157	1.00 20.68
ATOM	680	c	GLU	90	16.701	17.229	48.645	1.00 25.59
ATOM	681	ō	GLU	90	15.833	18.042	48.368	1.00 21.57
								1.00 21.37
MOTA	682	CB	GLU	90	16.031	15.816	50.682	1.00 22.21
ATOM	683	CG	GLU	90	15.782	14.403	51.228	1.00 37.59
ATOM	684	CD	GLU	90	17.071	13.641	51.447	1.00 83.49
ATOM	685	OE1	GLU	90	18.179	14.151	51.342	1.00 54.20
ATOM	686	OE2	GLU	90	16.875	12.373	51.749	1.00 64.65
ATOM	687	N	GLY	91	17.977	17.509	48.510	1.00 21.39
ATOM	688	CA	GLY	91	18.394	18.769	47.906	1.00 17.77
ATCM	689			91	18.673		48.839	
		C	CLY			19.911		
MOTA	690	0	GLY	91	18.769	19.764	50.055	1.00 16.81
MOTA	691	И	TYR	92	18.861	21.086	48.225	1.00 13.02
ATOM	692	CA	TYR	92	19.143	22.266	48.994	1.00 10.33
ATOM	693	C	TYR	92	18.575	23.478	48.347	1.00 9.87
ATOM	694	0	TYR	92	18.270	23.483	47.144	1.00 15.89
MOTA	695	CB	TYR	92	20.678	22.488	49.278	1.00 15.40
MOTA	696	CG	TYR	92	21.546	22.468	48.012	1.00 15.13
ATOM	697	CD1	TYR	92	21.620	23.576	47.166	1.00 14.75
ATOM	698	CD2	TYR	92	22.317	21.350	47.683	1.00 16.09
HOTA	699	CE1	TYR	92	22.404	23.561	46.008	1.00 6.50
ATOM	700	CE2	TYR	92	23.067	21.300	46.504	1.00 15.12 1.00 18.13 1.00 13.37
ATOM	701	CZ	TYR	92	23.156	22.424	45.683	1.00 12.13
ATOM	702	OH	TYR	92	23.944	22.393	44.517	1.00 13.37
ATOM	703	N	VAL	93	18.447	24.504	49.189	1.00 11.93
ATOM	704	CA	VAL	93	18.025	25.822	48.778	1.00 14.74
ATOM	705	c	VAL	93	19.281	26.666		1.00 16.00
							48.625	00 16.00
ATOM	706	0	VAL	93	20.172	26.625	49.451	1.00 15.16
ATOM	707	CB	VAL	93	17.073	26.480	49.791	1.00 23.45
ATOM	708	CG1	VAL	93	16.855	27.937	49.413	1.00 26.05
ATOM	709	CG2	VAL	93	15.716	25.764	49.771	1.00 22.90
ATOM	710	N	GLN	94	19.361	27.345	47.521	1.00 13.78
ATOM	711	CA	GLN	94	20.480	28.195	47.227	1.00 10.53
ATOM	712	c	GLN	94	19.948	29.583	46.998	1.00 12.23
ATOM	713	ŏ	GLN	94	19.153	29.788	46.061	1.00 15.52
ATOM	714	СВ	GLN	94	21.232	27.727		
							45.934	1.00 7.95
ATOM	715	CG	GLN	94	22.361	28.708	45.469	1.00 11.37
MOTA	716	CD	GLN	94	23.431	27.999	44.632	1.00 12.04
ATOM	717		GLN	94	23.805	26.879	44.946	1.00 13.60
ATOM	718	NE2	GLN	94	23.719	28.527	43.449	1.00 7.98
ATOM	719	M	,CLU	95	20.396	20.531	47.820	1.00 11.78
ATOM	720	CA	GLU	95	19.974	31.899	47.643	1 00 17 47
MOTA	721	c.	GLU	95	21.149	32.804	47.398	1.00 13.47
ATOM	722	Ö	GLU	95				00 15.42
ATOM	723	СВ			22.206	32.623	47.985	19.23
			GLU	95	19.277	32.427	48.878	1.00 12.52
ATOM	724	CG	GLU	95	18.009	31.684	49.215	1.00 22.46
MOTA	725	CD	GLU	95	17.657	32.016	50.622	1.00 45.93
ATOM	726	OEL	GLU	25	17.574	33.166	51.011	1.00100.00
MOTA	727	OE2	GLU	9,5	17.764	30.987	51.423	1.00100.00
ATOM	728	37	ARG	96	20.929	33.838	46.601	1.00 15.31
ATOM	729	CA	ARG	7.6	11.978	34.783	46.342	1.00 16.87
ATOM	730	c	ARG	76	11.510	36.195		
A10	, 30	-	ANG	70		10.112	46.206	1.00 15.84

ATOM	731	0	ARG	96	20.389	36.488	45.806	1.00 15.01
ATOM	732	CB.	ARG	96	22.582	34.463	14.967	1.00 16 19
ATOM	733	CG	ARG	96	23.495	33.247	44.929	1.00 17.61
ATOM	734	CD	ARG	96	24.615	33.453	43.908	1.00 9.06
ATOM	735	ΝE	ARG	96	25.411	32.277	43.766	1.00 9.88
ATOM	736	CZ	ARG	96	25.434	31.493	42.693	1.00 20.03
ATOM	737	SHI	ARG	96	24.684	31.709	41.615	1.00 15.29
ATOM	738	::H2	ARG	96	26.236	30.430	42.714	1.00 11.03
ATOM	739	::	THR	97	22.470	37.068	46.344	1.00 13.39
ATOM	740	CA	THR	97	22.368	38.424	45.935	1.00 13.12
ATOM	741	C	THR	97	23.593	38.688	45.084	1.00 16.81
ATOM	742	0	THR	97	24.686	38.347	45.485	1.00 19.25
ATOM	743	CB	THR	97	22.282	39.442	47.066	1.00 26.27
ATOM	744	OG1	THR	97	21.225	39.101	47.945	1.00 31.43
ATOM	745	CG2	THR	97	22.038	40.804	46.445	1.00 15.90
ATOM	746	N	ILE	98	23.396	39.219	43.899	1.00 16.23
ATOM	747	CA	ILE	98	24.486	39.526	42.977	1.00 16.70
ATOM	748	С	ILE	98	24.533	41.017	42.686	1.00 21.10
ATOM	749	0	ILE	98	23.628	41.566	42.075	1.00 14.58
ATOM	750	CB	ILE	98	24.385	38.752	41.660	1.00 13.47
ATOM	751	CG1	ILE	98	24.480	37.236	41.890	1.00 16.09
ATOM	752	CG2	ILE	98	25.457	39.231	40.679	1.00 13.30
MOTA	753	CD1	ILE	98	23.875	36.431	40.738	1.00 13.93
ATOM	754	31	PHE	99	25.613	41.678	43.110	1.00 14.86
ATOM	755	CA	PHE	99	25.719	43.098	42.896	1.00 12.44
ATOM	756	C	PHE	9.9	26.514	43.441	41.699	1.00 20.37
ATOM	757	0	PHE	99	27.696	43.164	41.700	1.00 20.07
ATOM	758	CB	PHE	99	26.401	43.770	44.084	1.00 15.96
ATOM	759	CG	PHE	99	25.638	43.624	45.356	1.00 21.41
ATOM	760	CD1	PHE	99	25.863	42.524	46.189	1.00 24.98
ATOM ATOM	761	CD2 CE1	PHE	99 99	24.698	44.585	45.743	1.00 22.94
ATOM	762 763	CE2	PHE	99	25.176 23.992	42.400	47.400	1.00 32.06
ATOM	764	CZ	PHE	99	24.235	44.469	47.771	1.00 24.26
ATOM	765	×	PHE	100	25.906	44.085	40.704	
ATOM	766	CA	PHE	100	26.679	44.522	39.554	1.00 12.53 1.00 8.75
ATOM	767	c	PHE	100	27.294	45.855	39.872	1.00 21.81
ATOM	768	õ	PHE	100	26.599	46.775	40.308	1.00 21.81 1.00 20.31 1.00 5.94
ATOM	769	C3	PHE	100	25.927	44.572	38.226	1.00 5.94
ATOM	770	CG	PHE	100	25.537	43.183	37.764	1.00 12.75
ATOM	771	CDI	PHE	100	24.426	42.538	38.325	1.00 16.31
ATOM	772	CD2	PHE	100	26.317	42.484	36.843	1.00 15.27
ATOM	773	CE1	PHE	100	24.087	41.230	37.975	1.00 13.50
ATOM	774	CE2	PHE	100	25.965	41.192	26.435	1.00 21.25
ATOM	775	CZ	PHE	100	24.852	40.567	37.014	1.00 21.06
ATOM	776	17	LYS	101	28.603	45.946	39.737	1.00 15.49
ATOM	777	CA	LYS	101	29.270	47.179	40.085	1.00 17.93
ATOM	778	С	LYS	101	28.732	48.349	39.287	1.00 13.71
MOTA	779	0	LYS	101	28.658	48.304	38.072	1.00 17.18
MOTA	780	CB	LYS	101	30.784	47.069	39.950	1.00 17.13
MOTA	781	CG	LYS	101	31.518	48.252	40.551	1.00 18.01
MOTA	782	CD	LYS	101	33.036	48.060	40.534	1.00 26.70
ATOM	783	CE	LYS	101	33.797	49.116	41.332	1.00 41.58
ATOM	784	37	ASP	:02	28.353	49.403	39.997	1.00 18.09
ATOM	785	CA	ASP	102	27.805	50.618	39.368	1.00 23.08
MOTA	786	C	`ASP	:02	26.559	50.356	38.549	1.00 25.42
ATOM	787	0	ASP	102	26.292	51.061	37.586	1.00 23.34
ATOM	788	CB	ASP	102	28.840	\$1.369	38.516	1.00 26.27
ATOM	789	CG	ASP	102	30.109	51.629 51.233	39.296	1.00 57.01
ATOM	790 791	001		102	31.206	11.233	38.931	1.00 63.33
MOTA	791	002		102	29.886	52.200	40.464	2.00 47.66
MOTA	793	:: CA	ASP	103	25.813	49.328	38.933	1.00 47.66 1.00 20.17 1.00 15.70 1.00 18.47
ATOM ATOM	793	CA	ASP	103	24.602	3.949	33.233	1.00 15.70
ATOM	795	ò	ASP	103	23.608	48.224	39.189	1.00 18.47
ATOM	796	C3	ASP	0.3	24.899	48.431	40.409	1.00 17.72
ATOM	797	CG	ASP	103	23.946	48.387	35.860	1.00 19.89
	,,,,		A.S.P	- 33	43.240	40.25/		00 23.93

ATOM	798	OD1	ASP	103	24.238	48.274	34.688	1.00 19.05
ATOM	799	OD2		103	22.774	48.809	36.283	1.00 23.89
ATOM	800	31	GLY	104	22.612	47.542	38.646	1.00 20.17
ATOM	801	CA	GLY	104	21.598	46.900	39.498	1.00 20.22
ATOM	802	C	GLY	104	22.055	45.619	40.180	1.00 24.68
ATOM	803	ŏ	GLY	104	23.202	45.211	40.085	1.00 18.06
ATOM	804	:1	ASN	105	21.125	44.967	40.872	1.00 15.71
ATOM	905	CA	ASN	105	21.425	43.703	41.510	1.00 8.89
ATOM	806	C	ASN	105	20.399	42.620	41.181	1.00 21.85
ATOM	807	ō	ASN	105	19.255	42.911	40.824	1.00 15.17
ATOM	808	CB	ASN	105	21.605	43.840	43.001	1.00 8.58
ATOM	809	CG	ASN	105	20.359	44.366	43.697	1.00 43.57
ATOM	810	001	ASN	105	19.565	43.601	44.259	1.00 36.67
ATOM	811	ND2	ASN	105	20.178	45.674	43.659	1.00 36.47
ATOM	812	N	TYR	106	20.826	41.365	41.328	1.00 16.80
ATOM	813	CA	TYR	106	19.966	40.219	41.156	1.00 13.90
ATOM	814	С	TYR	106	19.763	39.543	42.475	1.00 11.05
ATOM	815	0	TYR	106	20.678	39.404	43.281	1.00 13.86
ATOM	816	CB	TYR	106	20.547	39.128	40.246	1.00 15.88
ATOM	817	CG	TYR	106	20.619	39.398	38.793	1.00 15.57
ATOM	818	CD1	TYR	106	19.952	40.458	38.178	1.00 13.14
ATOM	819	CD2	TYR	106	21.373	38.524	38.006	1.00 13.35
MOTA	820	CE1	TYR	106	20.038	40.632	36.793	1.00 13.44
ATOM	821	CE2	TYR	106	21.481	38.692	36.628	1.00 10.87
ATOM	822	CZ	TYR	106	20.814	39.751	36.025	1.00 15.93
ATOM	823	ОН	TYR	106	20.970	39.931	34.670	1.00 17.32
ATOM	824	21	LYS	107	18.538	39.115	42.709	1.00 12.39
ATOM	825	CA	LYS	107	18.194	39.349	43.897	1.00 11.51
ATOM	826	С	LYS	107	17.619	37.037	43.397	1.00 17.25
ATOM	827	0	LYS	107	16.704	37.010	42.562	1.00 13.14
ATOM	828	CB	LYS	107	17.217	39.063	44.823	1.00 14.82
ATOM	829	CG	LYS	107	17.860	39.631	46.060	1.00 40.71
ATOM	830	CD	LYS	107	18.528	40.974	45.793	1.00 43.48
ATOM	831	N	THR	108	18.205	35.951	43.835	1.00 14.95
ATOM	832	CA	THR	108	17.774	34.658	43.352	1.00 11.97
ATOM	833	С	THR	108	17.463	33.696	44.468	1.00 15.81
ATOM	834	0	THR	108	18.043	33.734	45.582	1.00 13.68
ATOM	835	CB	THR	108	18.847	34.034	42.410	1.00 23.81
ATOM	836	OGl	THR	108	20.064	33.791	43.137	1.00 13.88
ATOM	837	CG2	THR	108	19.123	34.968	41.264	1.00 13.04
MOTA	838	::	ARG	109	16.560	32.804	44.154	1.00 13.57
MOTA	839	CA	ARG	109	16.212	31.751	45.048	1.00 12.56
ATOM	840	C	ARG	109	15.939	30.498	44.254	1.00 13.07
MOTA	841 842	0	ARG	109	15.239	30.509	43.249	1.00 12.52
ATOM	843	CB	ARG	109	15.069	32.100	45.959	1.00 17.32
ATOM ATOM	844	CG	ARG	109	14.767	30.995 31.160	46.932 47.610	1.00 17.92
ATOM	845	NE	ARG	109	12.821	29.854	47.883	
ATOM	846	CZ	ARG	109	12.968	29.244	49.035	1.00 36.05
ATOM	847	NH1		109	13.630	29.815	50.046	1.00 44.11
ATOM	848	NH2	ARG	109	12.432	28.041	49.195	1.00 94.34
ATOM	849	11	ALA	110	16.577	29.414	44.635	1.00 13.26
ATOM	850	CA	ALA	. 10	16.377	28.207	43.870	1.00 12.68
ATOM	851	c	ALA	110	16.346	26.979	44.734	1.00 13.15
ATOM	852	0	ALA	110	16.829	26.965	45.869	1.00 16.75
ATOM	853	CB	'ALA	110	17.465	28.059	42.822	1.00 17.31
ATOM	354	3	GLU	:11	15.770	25.939	44.176	1.00 15.39
ATOM	855	CA	GLU	111	15.741	24.655	44.823	1.00 15.24
ATOM	856	c	GLU	111	16.438	23.678	43.926	1.00 12.08
ATOM	857	ō	GLU	111	16.086	23.545	42.771	1.00 15.70
ATOM	858	CB	GLU	111	14.303	24.122	44.993	
ATOM	359	CG	GLU	111	13.744	24.242	46.399	1.00 19.20
ATOM	360	CD	GLU	.11	12.247	24.280	46.372	1.00 60.99
ATOM	861	OE1	GLU		11.589	23.843	45.432	1.00 76.05
ATOM	362	9E2		111	11.742	24.956	47.280	1.00 54.87
MOTA	363	12	TAL	112	17.438	22.965	44.457	1.00 10.73
MOTA	364	ÇA	TAL	112	18.063	11.978	43.631	1.00 10.98

ATOM	865	С	VAL	112	17.968	20.630	44.261	1.00 8.62
ATOM	866	ō	VAL	112	18.271	20.438	45.432	1.00 8.62
HOTA	867	СВ	VAL	112	19.428	22.358	43.012	1.00 22.75
ATOM	868	CG1	VAL	112	19.966	23.704	43.487	1.00 16.69
ATOM	869	CG2	VAL	112	20.452	21.232	43.078	1.00 18.47
ATOM	870	N	LYS	113	17.415	19.732	43.516	1.00 14.67
ATOM	871	CA	LYS	113	17.175	18.421	44.045	1.00 16.41
ATOM	872	С	LYS	113	16.822	17.485	42.931	1.00 16.41
MOTA	873	ò	LYS	113	16.695	17.893	41.808	1.00 16 27
MOTA	874	CB	LYS	113	16.032	18.497	45.036	1.00 22.50
MOTA	875	CG	LYS	113	14.792	19.084	44.376	1.00 20.40
MOTA	876	CD	LYS	113	13.509	18.321	44.703	1.00 44.65
ATOM	877	CE	LYS	113	12.526	19.134	45.528	1.00 54.02
MOTA	878	NZ	LYS	113	12.379	20,518	45.036	1.00100.00
ATOM	879	N	PHE	114	16.683	16.208	43.267	1.00 10.09
ATOM	880	CA	PHE	114	16.325	15.175	42.317	1.00 11.41
ATOM	881	С	PHE	114	14.806	14.975	42.181	1.00 14.18
ATOM	882	0	PHE	114	14.110	14.878	43.160	1.00 15.03
ATOM	883	CB	PHE	114	16.866	13.838	42.838	1.00 12.89
ATOM	884	CG	PHE	114	18.231	13.536	42.338	1.00 16.80
ATOM	885	CD1	PHE	114	19.344	13.795	43.139	1.00 18.61
ATOM	886	CD2	PHE	114	18.403	13.009	41.056	1.00 19.50
MOTA	887	CEL	PHE	114	20.627	13.500	42.665	2.00 22.78
MOTA	888	CE2	PHE	114	19.673	12.708	40.572	1.00 25.36
ATOM	889	CZ	PHE	114	20.780	12.953	41.387	1.00 25.36
ATOM	890	14	GLU	:15	14.354	14.819	40.966	. 00 12 20
MOTA	891	CA	GLU	115	12.978	14.473	40.642	1.00 11.40 1.00 13.30 1.00 18.72
MOTA	892	C	GLU	115	13.121	13.193	39.906	1.00 13.30
ATOM	893	0	GLU	115	13.434	13.207	38.730	1.00 18.72
ATOM	894	CB	GLU	115	12.348	15.481	39.667	1.00 9.68
ATOM	895	CG	GLU	115	11.856	16.747	40.376	1.00 19.54
ATOM	896	CD	GLU	115	10.742	16.460	41.342	1.00 38.12
ATOM	897	OE1	GLU	115	10.181	15.395	41.431	1.00 34.84
ATOM	898	OE2	GLU	115	10.460	17.461	42.079	1.00 27.88
MOTA	899	N	GLY	116	13.005	12.087	40.585	1.00 14.51
ATOM	900	CA	GLY	116	13.225	10.861	39.869	1.00 15.91
MOTA	901	С	GLY	116	14.727	10.767	39.641	1.00 23.59
MOTA	902	0	GLY	116	15.516	10.922	40.570	1.00 19.35
MOTA	903 904	N CA	ASP	117	15.137	10.564	38.439	00 20.20
MOTA MOTA	904	CA	ASP ASP	117	16.572	10.462	38.233	1.00 28.00
ATOM	906	ŏ	λSP	117	17.237 18.423	11.677	37.598	1.00 22.39
		СВ				11.672	37.265	1.00 21.33
ATOM ATOM	907 908	CG	ASP	117	17.055	9.074	37.733	1.00 33.06 1.00 55.04 1.00 59.57
ATOM	909	ODI	ASP		16.624	8.677	36.348	1.00 55.04
ATOM	910	OD2	ASP	117	16.230	9.468	35.495	1.00 59.57
ATOM	911	N N	THR	118	16.805 16.463	7.391	36.130	1.00 82.48
ATOM	912	CA	THR	118	16.889	12.729	37.493 36.910	1.00 19.62
MOTA	913	c	THR	118	17.186	14.988	37.976	1.00 18.21
ATOM	914	ŏ	THR	118 118	16.498	15.064	38.996	1.00 18.92
ATOM	915	CB	THR	118	15.806	14.497	35.952	1.00 19.03
ATOM	916	OG1	THR	118	15.552	13.508	34.990	1.00 21.42
ATOM	917	CG2	THR	118	16.217	15.793	35.275	1.00 21.42
ATOM	918	N	LEU	- 10	18.284	15.681	37.805	1.00 13.66
MOTA	919	0.5	* ****	119 119 119	18.679	16.706	38.759	1.00 13.50
ATOM	920	c	LEU	119	18.036	17.992	38.269	1.00 13.50
ATOM	921	ō	LEU	119	18.194	18.368	37.091	.00 12.49
ATOM	922	СВ	LEU	: 19	20.243	15.815	38.839	1.00 12.25
ATOM	923	CG	LEU	119	20.845	17.678	39.951	
ATOM	924	CDI	LEU	. 10	20.701	19.167	39.669	1.00 10.11
ATOM	925	CD2	LEU	119	20.366	17.311	41.333	1.00 3.90 1.00 10.11 1.00 7.36
ATOM	926	N	VAL	20	17.230	13.595	39.170	1.30 13.34
ATOM	927	CA	'/AL	120	16.466	. 9 797	38.859	1.00 13.77
ATOM	928	c.	VAL	120	16.929	21.039	39.587	1.00 8.56
ATOM	929	0	VAL	120	17.135	01.039	40.752	1.00 13.32
ATOM	930	C3	VAL	120	14.939	19.566	39.082	1.00 17.50
ATOM	931	CG1	VAL	120	14.133	20.790	33.642	1.00 17.58

ATOM	932	CG2	VAL	120	14.501	18.351	38.246	1.00 15.35
ATOM	933	11	ASN	121	17.067	22.111	38.839	1.00 15.35
ATOM	934	CA	ASN	121	17.424	23.405	39.400	1.00 11.78
								1.00 11.78
ATOM	935	C	ASN	121	16.301	24.382	39.060	1.00 11.18
ATOM	936	0	ASN	121	16.195	24.802	37.934	1.00 11.09
ATOM	937	CB	ASN	121	18.753	23.928	38.791	1.00 11.41
ATOM	938	CG	ASN	121	19.201	25.261	39.367	1.00 11.07
ATOM	939	OD 1	ASN	121	18.773	25.654	40.461	1.00 12.06
MOTA	940	ND2	ASN	121	20.124	25.938	38.670	1.00 12.06
ATOM	941	N	ARG	122	15.470	24.706	40.029	1.00 13.69
ATOM	942	CA	ARG		14.348	25.610	39.825	1.00 12.99
ATOM	943	C		122				
			ARG	122	14.622	26.946	40.498	1.00 5.89
MOTA	944	0	ARG	122	14.749	27.011	41.723	1.00 14.47
ATOM	945	CB	ARG	122	13.068	25.025	40.417	1.00 15.99
ATOM	946	CG	ARG	122	12.478	23.921	39.589	1.00 30.23
ATOM	947	CD	ARG	122	11.282	23.244	40.281	1.00 60.61
ATOM	948	N	ILE	123	14.663	27.992	39.680	1.00 11.46
ATOM	949	CA	ILE	123	15.030	29.340	40.095	1.00 11.86
ATOM	950	C	ILE	123	13.991	30.450	39.835	1.00 10.54
ATOM	951	ö	ILE	123	13.370	30.535	38.765	1.00 12.83
ATOM	952	CB		123	16.296	29.757	39.292	1.00 12.83
			ILE					1.00 15.41
ATOM	953	CG1	ILE	123	17.316	28.585	39.180	1.00 12.27
ATOM	954	CG2	ILE	123	16.944	30.993	39.918	1.00 14.01
MOTA	955	CD1	ILE	123	17.652	28.242	37.743	1.00 7.74
ATOM	956	N	GLU	124	13.953	31.358	40.793	1.00 11.36
MOTA	957	CA	GLU	124	13.189	32.572	40.700	1.00 15.20
MOTA	958	Ċ	GLU	124	14.168	33.713	40.811	1.00 11.93
ATOM	959	ō	GLU	124	14.919	33.797	41.780	1.00 15.61
ATOM	960	CB	GLU	124	12.028	32.677	41.751	1.00 19.74
ATOM	961	CG	GLU	124	12.387	33.337	43.089	1.00 72.94
ATOM	962	N	LEU	125	14.183	34.550	39.808	
MOTA	963	CA	LEU	125	15.092	35.654	39.767	1.00 15.00
ATOM	964	С	LEU	125	14.420	37.011	39.722	1.00 19.35
MOTA	965	0	LEU	125	13.563	37.267	38.893	1.00 18.41
ATOM	966	CB	LEU	125	15.976	35.533	38.510	1.00 14.29
ATOM	967	CG	LEU	125	17.003	36.683	38.375	1.00 17.55
MOTA	968	CD1	LEU	125	18.302	36.083	37.849	1.00 13.46
ATOM	969	CD2	LEU	125	16.511	37.732	37.367	1.00 12.09
ATOM	970	N	LYS	126	14.890	37.897	40.554	
ATOM	971	CA	LYS	126	14.391	39.260	40.579	1.00 15.92
ATOM	972	C	LYS	126	15.563	40.276	40.445	1.00 12.73 1.00 15.92 1.00 18.53 1.00 19.86 1.00 17.31 1.00 33.94
	973							1.00 18.53
ATOM		0	LYS	126	16.489	40.246	41.246	1.00 19.86
ATOM	974	CB	LYS	126	13.611	39.487	41.877	1.00 17.31
MOTA	975	CG	LYS	126	12.853	40.786	41.923	1.00 33.94
ATOM	976	CD	LYS	126	11.366	40.601	41.675	1.00 60.87
ATOM	977	CE	LYS	126	10.652	41.929	41.521	1.00 52.70
ATOM	978	NZ	LYS	126	11.229	42.988	42.367	1.00 47.22
ATOM	979	N -	GLY	127	15.514	41.127	39.411	1.00 18.71
ATOM	980	CA	GLY	127	16.551	42.151	39.121	1.00 17.32
ATOM	981	C	GLY	127	16.012	43.572	39.272	1.00 25.32
· ATOM	982	õ	GLY	127	14.981	43.908	38.693	1.00 20.14
ATOM	983	N	ILE	128				. 00 10 40
					16.706	44.404	40.070	1.00 18.42
ATOM	984	CA	ILE	128	16.282	45.787	40.243	1.00 21.04
MOTA	985	С	ILE	128	17.405	46.789	40.196	1.00 21.04
ATOM	986	0	ILE	128	18.562	46.496	40.429	1.00 19.37
ATOM	987	CB	ILE	128	15.482	46.052	41.504	1.00 23.82
ATOM	988	CGI	ILE	128	16.408	45.888	42.701	1.00 23.26
ATOM	989	CG2	ILE	128	14.272	45.120	41.577	1.00 28.95
ATOM	990	CD1	ILE	128	15.824	46.391	44.013	
ATOM	991	N	ASP	129	15.824 16.999 17.861 18.864	-8.002	39.918	1.00 29.89
ATOM	992	CA	ASP	29	7.961	49.124	39.882	1.00 13.53
ATOM	393	C	ASP	: 29	. 0 0 . 1	19.086	38.801	1.00 18.53
				129	.3.864			20.35
ATOM	994	0	ASP	. 29	19.949	49.632	38.953	
MOTA	995	CB	ASP	129	13.498	49.407	41.253	1.00 20-57
ATOM	996	CG	ASP	129	17.545	50.077	42.226	1.00 20.57
MOTA	997	001	ASP	129 129	16.653	50.842	41.883	1.00 49.42
ATOM	998	OD2	ASP	129	17,770	49.740	43.475	1.00 38.07
		_				_	_	

MOTA	999	22	PHE	130	18.510	48.493	37.693	1.00 16.40
ATOM	1000	Cλ	PHE	130	19.433	48.459	36.563	1.00 16.99
ATOM	1001	c	PHE	130	19.330	49.732	35.756	1.00 35.37
ATOM	1002	õ	PHE	130	18.242	50.318	35.623	1.00 27.34
ATOM	1003	СЗ	PHE	130	19.248	47.223	35.657	1.00 18.07
ATOM	1004	CG	PHE	130	19.809	45.980	36.312	1.00 19.10
ATOM	1005	CDI	PHE	130	19.021	45.210	37.171	1.00 16.15
ATOM	1006	CD2	PHE	130	21.126	45.572	36.073	1.00 19.17
ATOM	1007	CEI	PHE	130	19.536	44.074	37.801	1.00 23.37
ATOM	1008	CE2	PHE	130	21.665	44.445	36.703	1.00 21.11
ATOM	1009	cz	PHE	130	20.867	43.703	37.575	1.00 22.13
ATOM	1010	3	LYS	131	20.464	50.169	35.218	1.00 31.09
ATOM	1011	CA	LYS	131	20.477	51.371	34.400	1.00 27.52
ATOM	1012	c	LYS	131	20.105	51.045	32.992	1.00 25.57
ATOM	1013	ò	LYS	131	20.695	50.169	32.343	1.00 22.97
ATOM	1014	СВ	LYS	131	21.796	52.109	34.438	1.00 32.64
ATOM	1015	CG	LYS	131	22.153	52.633	35.813	1.00 38.34
ATOM	1016	CD	LYS	131	23.646	52.886	35.975	1.00 75.76
ATOM	1017	3	GLU	132	19.116	51.751	32.509	1.00 26.88
ATOM	1018	CA	GLU	132	18.623	51.484	31.189	1.00 28.42
ATOM	1019	C	GLU	132	19.710	51.514	30.140	1.00 36.19
ATOM	1020	ŏ	GLU	132	19.617	50.862	29.101	1.00 39.24
ATOM	1021	CB.	GLU	132	17.374	52.331	30.830	1.00 29.04
ATOM	1021 1022	7	ASP	133	20.752	52.254	30.438	1.00 40.08
ATOM	1023	CA	ASP	133	21.883	52.442	29.525	1.00 45.36
MOTA	1024	č	ASP	133	23.224	51.861	30.049	1.00 50.61
MOTA	1025	ő	ASP	133	24.299	52.243	29.572	1.00 52.14
ATOM	1025	CЗ	ASP	133	22.063	53.946	29.332	1.00 50.45
MOTA	1027	CG	ASP	133	22.109	54.642	30.670	1.00 87.10
MOTA	1028	OD1	ASP	133	21.408	54.314	31.624	1.00 91.27
ATOM	1029	OD2	ASP	133	23.047	55.552	30.739	1.00100.00
MOTA	1030	14	GLY	134	23.159	50.970	31.053	1.00 37.06
MOTA	1031	CA	GLY	134	24.349	50.376	31.639	1.00 30.22
ATOM	1032	C	GLY	134	24.845	49.228	30.803	1.00 23.10
MOTA	1033	0	GLY	134	24.360	48.990	29.685	1.00 19.23
ATOM	1034	34	ASN	135	25.807	48.486	31.341	1.00 18.66
ATOM	1035	CA	ASN	1.35	25.239	47.370	30.563	1.00 18.03
ATOM	1036	C	ASN	135	25.372	46.199	30.406	1.00 15.75
MOTA	1037	CB	ASN	135	25.485	45.430	29.461	1.00 16.03
MOTA	1039	CG	ASN	135	27.665 28.743	46.883	31.139	1.00 19.27
ATOM ATOM	1040	OD1	ASN	135	28.969	47.943 48.595	30.078	1.00 15.75 1.00 16.03 1.00 19.27 1.00 20.99 1.00 25.69
ATOM	1041	:ID2	ASN	135	29.423	48.095	32.239	1.00 22.57
ATOM	1042	31	ILE	136	24.444	46.052	31.362	1.00 18.14
ATOM	1043	CA	ILE	136	23.494	44.924	31.368	1 00 19 78
MOTA	2044	c	ILE	136	22.331	45.086	30.384	1.00 23.76
ATOM	1045	ō	ILE	136	22.178	44.313	29.395	1.00 22.53
ATOM	1046	CB	ILE	136	23.078	44.500	32.804	1.00 23.76 1.00 22.53 1.00 21.24
ATOM	1047	CG1	ILE	136	24.230	43.728	33.423	1.00 28.44
ATOM	1048	CG2	ILE	136	21.899	43.543	32.770	1.00 22.77
MOTA	1049	CD1	ILE	136	25.346	44.596	33.935	1.00 12.29
· ATOM	1050	24	LEU	137	21.543	46.117	30.640	1.00 18.21
ATOM	2051	CA	LEU	137 137 137	20.394	46.415	29.815	1.00 23.30
MOTA	1052	С	LEU	137	20.828	46.875	28.470	1.00 27.26
ATOM	1053	0	LEU	137	20.181	46.619	27.488	1.00 27.00
ATOM	1054		, CEA	137	19.442	47.430	30.490	1.00 21.74
MOTA	1055	CG	LEU	_37	18.828	46.852	31.762	1.00 27.00 1.00 21.74 1.00 22.56
ATOM	1056		LEU	137	17.856	47.837	32.415	1.00 22.27
ATOM	1057	CD2	LEU	137	18.119	45.554	31.424	1.00 37.52
ATOM	1058	3	GLY	138	21.979	47.527	28.432	1.00 22.14
ATOM	:059	CA	GLY	138	22.510	48.033	27.187	1.00 20.03
ATOM	1060	5	GLY	138	23.157	46.959	26.368	1.00 20.16
ATOM ATOM	1062	0	GLY	138	23.600	47.202	25.264	1.00 22.44
ATOM	1063	ÇA.	HIS		23.246	44.655	26.903 26.148	00 19.27
ATOM	1064		HIS	139	25.301	44.929	25.148	1.00 20.24
ATOM	1065	5	HIS	:39	25,605	44.745	14.439	1.00 20.13
A+0.1	2000	-			40.000	-4.743	.4.433	1.00 17.37

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ATOM	1066	СВ	HIS	139	22.931	44.207	25.018	1.00 22.20
ATOM	1067	CG	HIS	:39	21.708	43.551	15.550	1.00 25.52
ATOM	1068	ND1	HIS	139	21.666	42.182	25.785	1.00 25.67
ATOM	1069		HIS	139	20.525	44.092	25.927	1.00 28.09
ATOM	1070		HIS	139	20.474	41.918	26.275	1.00 27.50
ATOM	1071	NE2	HIS	139	19.766	13.044	25.382	1.00 29.53
ATOM	1072	N	LYS	140	26.187	45.311	26.525	1.00 23.51
ATOM	1073	CA	LYS	140	27.569	45.638	26.197	1.00 25.82
ATOM	1074	c	LYS	140	28.600	44.537	26.560	1.00 25.82
ATOM	1075	o	LYS	140	29.824	44.730	26.391	1.00 22.29
ATOM	1076	CB	LYS	140	27.977	46.937	26.911	1.00 27.56
ATOM	1077	CC	LYS	140	27.269	48.217	26.445	1.00 31.19
ATOM	1078	CD	LYS	140	27.234	49.254	27.582	1.00 51.32
ATOM	1079	CE	LYS	140	26.924	50.696	27.169	1.00 47.92
ATOM	1080	NZ	LYS	140	27.112	51.663	28.284	1.00 73.76
ATOM	1081	N	LEU	141	28.116	43.403	27.115	1.00 19.33
ATOM	1082	CA	LEU	141	28.987	42.296	27.559	1.00 14.32
MOTA	1083	С	LEU	141	29.366	41.401	26.427	1.00 20.75
ATOM	1084	0	LEU	141	28.526	41.087	25.620	1.00 19.01
MOTA	1085	CB	LEU	141	28.313	41.488	28.676	1.00 12.53
MOTA	1086	CG	LZU	141	27.979	42.352	29.875	1.00 17.54
ATOM	1087	CD1	LEU	141	27.700	41.469	31.070	1.00 24.81
ATOM	1088	CD2	LEU	141	29.116	43.310	30.182	1.00 27.50
MOTA	1089	И	GLU	142	30.644	40.987	26.346	
ATOM ATOM	1090	CA	GLU	142	31.040	40.059 38.691	25.311	1.00 13.43
ATOM	1091	0	GLU	142	30.442	38.393	26.787	1.00 16.43
ATOM	1093	CB	GLU	142	32.558	39.866	25.204	1.00 14.73
ATOM	1094	CG	GLU	142	33.290	41.077	24.624	1.00 29.30
ATOM	1095	CD	GLU	142	34.787	41.003	24.825	1.00 56.32
ATOM	1096	OE1	GLU	142	35.340	40.098	25.420	1.00 31.70
ATOM	1097	OE2	GLU	142	35.430	42.015	24.321	1.00 34.10
ATOM	1098	N	TYR	143	30.365	37.873	24.632	1.00 16.30
ATOM	1099	CA	TYR	143 143	29.837	36.542	24.764	1.00 20.04
ATOM	1100	С	TYR	143	30.925	35.559	25.049	1.00 12.46
ATOM	1101	0	TYR	143	31.327	34.792	24.193	1.00 16.99
MOTA	1102	СB	TYR	143	29.035	35.113	23.498	1.00 20.96
ATOM	1103	CG	TYR	143	28.187	34.857	23.674	1.00 16.12
ATOM	1104	CD1	TYR	143 143 143	27.040	34.859	24.472	1.00 18.24
ATOM	1105	CD2	TYR		28.512	33.684	12.986	1.00 12.87
ATOM	1106	CEI	TYR	143	26.257	33.708	24.615	1.00 17.91
ATOM	1107	CE2	TYR	143	27.735	32.530	23.104	1.00 16.58
ATOM ATOM	1108	CZ	TYR	:43 :43	26.603 25.861	31.432	23.914	1.00 17.35
ATOM	1110	OH N	ASN	144	31.392	35.597	26.251	1.00 23.40 1.00 12.40
ATOM	1111	CA	ASN	144	32.428	34.703	25.689	1.00 12.05
ATOM	1112	C	ASN	:44	32.433	34.675	28.193	1.00 15.75
ATOM	1113	õ	ASN		31.637	35.369	28.837	1.00 14.58
ATOM	1114	CB	ASN	- 44	33.823	35.038	25.068	1.00 18.45
ATOM	1115	CG	ASN	144 144 144 144 145	34.310	36.445	25.374	1.00 18.98
ATOM	1116	OD1	ASN	.44	34.150	36.951	27.488	1.00 20.34
ATOM	1117	ND2	ASN	144	34.891	37.085	23.382	1.00 23.02
MOTA	:118	N	TYR	145	33.311	33.876	23.773	1.00 12.16
A TOM	:119	CA	TYR	.45	33.343	33.765	30.195	1.00 10.63
ATOM	1120	C	TYR	145	34.765	33.458	30.730	1.00 14.58
ATOM	1121	0	TYR.	145	35.510	32.751	30.090	1.00 18.83
ATOM	1122	CB	TYR	145	32.404	32.627	30.571 11.826 11.808 13.030	1.00 9.76
ATOM	1123	CC	TYR	-45	31.698	12.916	1.826	1.00 11.86
ATOM	1124	CD1	TYR	.45	30.515	33.658	808	1.00 9.04
ATOM	1125	CD2	TYR	145	32.188	32.419	030	1.00 10.07
ATOM ATOM	1126	CE1	TYR	145	29.860	33.948	12.999	1.00 8.36 1.00 15.32
ATOM	1120	CEZ	TYR	145 145	31.544 30.375	32.707	34.235 34.206 35.376	1.00 15.32
ATOM	1128	OH	TYR	145	29.730	13.735	3 3 3 7 5	1.00 11.69
ATOM	1130	31	ASN	16	35.086	33.931	31.933	1.00 15.23
ATOM	1131	CA	ASN	146	36.415	33.737	32.550	1.00 15.36
ATOM	1132	C	ASN	146	36.426	32.518	33.589	1.00 19.68
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ATOM	1133	0	ASN	146	35.395	32.043	33.848	1.00 14.71
ATOM	1134	CB	ASN	146	36.844	35.062	33.235	1.00 11.89
ATOM	1135	CG	ASN	. 46	37.013	36.147	32.215	1.00 35.45
ATOM	1136	ODI	ASN	146 146	37.533	35.890	31.105	1.00 31.63
ATOM	1137	ND2	ASN	146	36.547	37.349	32.553	1.00 19.74
ATOM	1138	N	SER	:47	37.630	32.338	34.201	1.00 12.09
	1139	CA		147			35.266	1.00 22.09
ATOM	1140	CA	SER	147	37.804 37.769	31.320	36.575	1.00 11.70
ATOM			SER	-47				
ATOM	1141	0_	SER	147 147	38.219	33.125	36.671	1.00 16.56
ATOM	1142	CB	SER	147	39.148	30.540	35.129	1.00 9.87
ATOM	1143	OG	SER	147	39.212	29.980	33.828	1.00 33.20
ATOM	1144	N	HIS	148	37.195	31.365	37.583	1.00 5.53
ATOM	1145	CA	HIS	148	37.090	31.998	38.850	1.00 8.06
ATOM	1146	C	HIS	148	37.346	31.038	39.949	1.00 11.30
ATOM	1147	0	HIS	148	37.328	29.844	39.754	1.00 16.87
ATOM	1148	CB	HIS	148	35.648	32.608	39.067	1.00 11.29
ATOM	1149	CG	HIS	148	35.215	33.554	37.972	1.00 10.84
MOTA	1150	NDl		148	34.548	33.121	36.836	1.00 12.77
ATOM	1151	CD2		148	35.403	34.887	37.851	1.00 8.82
ATOM	1152	CE1	HIS	:48	34.389	34.178	36.060	1.00 8.84
ATOM	1153	NE2	HIS	148	34.882	35.242	36.647	1.00 8.82
ATOM	1154	N	ASN	:49	37.534	31.579	41.125	1.00 10.80
ATOM	1155	CA	ASN	149	37.626	30.805	42.345	1.00 13.35
ATOM	1156	С	ASN	:49	36.409	31.157	43.205	1.00 14.47
ATOM	1157	ō	ASN	149	36.099	32.320	43.387	1.00 18.17
ATOM	1158	CB	ASN	149	38.890	31.093	43.184	1.00 12.67
ATOM	1159	CG	ASN	149	40.148	30.822	42.424	1.00 20.21
ATOM	1160	OD1	ASN	:49	40.993	31.713	42.281	1.00 56.34
ATOM	1161	ND2	ASN	:49	40.210	29.641	41.818	1.00 16.44
ATOM	1162	N	VAL	150	35.773	30.144	43.741	1.00 14.65
ATOM	1163	CA	VAL	150	34-588	30.262	44.552	1.00 12.92
ATOM	1164	c	VAL	150	34.910	29.806	45.943	1.00 16.30
ATOM	1164	ŏ	VAL	150	35.257	28.665	46.147	1.00 17.83
ATOM	1166	CB	VAL	150	33.482	29.382	43.914	1.00 15.22
ATOM	1167	CG1		150	32.252	29.297	44.765	1.00 14.09
ATOM	1168		VAL	150	33.172	29.791	42.464	1.00 10.94
ATOM	1169	N N	TYR	151	34.796	30.716	46.900	1.00 17.64
ATOM	1170	CA	TYR	151	35.139	30.440	48.275	1.00 18.31
ATOM	1171	c	TYR	151	34.003	29.917	49.117	1.00 24.35
ATOM	1172	õ	TYR	151	32.963	30.536	49.239	1.00 20.83
ATOM	1173	СВ	TYR	151	35.793	31.681	48.920	1.00 20.15
ATOM	1174	CG	TYR	151	37.025	32.033	48.141	1.00 25.86
ATOM	1175		TYR	151	37.003	32.989	47.127	1.00 26.00
ATOM	1176	CD2		151	38.200	31.315	48.355	1.00 28.66
ATOM	1177		TYR	151	38.151	33.234	46.369	1.00 33.73
	1178			151	39.360	31.550		
ATOM	1179	CE2	TYR	151	39.325	32.512	47.619	
MOTA	1180		TYR	151	40.449	32.737	46.618	
ATOM ATOM		N	ILE	:52	34.250	28.791	45.877	
	1181	CA		152	33.255	28.159		
ATOM		CA	ILE	-32			50.572	1.00 14.12
ATOM	1183		ILE	152 152	33.619	28.056	52.000	1.00 18.51
ATOM	1184	O CB		152	34.728	27.703	52.336	1.00 22.05
ATOM	1185		ILE	152	32.979	26.776	50.060	1.00 16.66
ATOM	1186	CG1	ILE	152	32.431	26.875	48.638	1.00 11.30
ATOM	1187	CG2	ILE	_52	32.017	26.078	51.021	1.00 17.96
ATOM	1188		LLE	152	32.377	25.559	47.949	1.00 13.48
ATOM	1189	И	MSE	153	32.623	28.278	52.841	1.00 17.41
ATOM	1190	CA	MSE		32.789	28.162	54.269	1.00 22.61
ATOM	1191	С	MSE	153	31.534	27.648	54.916	1.00 27.31
ATOM	1192	0	MSE	153	30.433	27.831	54.396	1.00 20.50
ATOM	1193	CB	MSE	153	33.145	29.490	54.855	1.00 19.11
ATOM	1194	CG	MSE	153	34.010	30.302	53.957	1.00100.00
ATOM	1195	SE	MSE	153	34.060	32.117	54.524	1.00100.00
ATOM	1196	CE	MSE	113 114	33.463	31.798	56.330	00 30.27
ATOM	:197	N	ALA	154	31.733	26.983	56.053	1.00 22.29
ATOM	1198	CA	ALA	154 154	30.669	26.389	56.796	1.00 22.66
ATOM	1199	C	ALA	154	29.820	27.401	57.552	1.00 29.00

ATOM	1200	0	ALA	154	30.274	28.457	57.960	1.00 27.02
ATOM	1201	СВ	ALA	154	31.224	25.336	57.744	1.00 19.78
ATOM					28.566	27.063	57.726	1.00 29.43
	1202	N	ASP	155				
ATOM	1203	CA	ASP	155	27.669	27.887	58.484	1.00 32.18
ATOM	204	С	ASP	155	26.976	27.019	59.511	1.00 44.51
ATOM	1204	0	ASP	155	25.898	26.492	59.274	1.00 39.56
ATOM	1206	CB	ASP	155	26.659	28.617	57.597	1.00 31.70
MOTA	_207	CG	ASP	155	26.140	29.851	58.247	1.00 49.89
ATOM	1208	OD1	ASP	155	26.595	30.297	59.277	1.00 46.67
ATOM	1209	OD2	ASP	155	25.187	30.422	57.565	1.00 76.07
ATOM	1210	N	LYS	156	27.646	26.816	60.629	1.00 46.37
ATOM	1211	CA	LYS	156	27.116	25.954	61.654	1.00 53.23
ATOM	1212		LYS	156	25.750	26.369	62.224	1.00 65.62
ATOM	-212	c				25.520	62.703	1.00 65.54
	1213	0	LYS	156	25.012	25.612	62.705	1.00 59.51
MOTA	1214	CB	LYS	156	28.147		62.725	
ATOM	1215	N	GLN	157	25.398	27.655	62.138	1.00 68.32
ATOM	.216	CA	GLN	157	24.119	28.135	62.670	1.00 73.00
ATOM	217	С	GLN	157	22.891	27.767	61.817	1.00 87.53
ATOM	1218	0	GLN	157	21.778	27.547	62.325	1.00 96.16
ATOM	1219	N	LYS	158	23.095	27.725	60.506	1.00 72.49
ATOM	:220	CA	LYS	158	22.040	27.386	59.593	1.00 66.19
ATOM	1221	C	LYS	158	22.235	25.985	59.040	1.00 58.21
ATOM	1221 1222 1223	ō	LYS	158	21.447	25.524	58.226	1.00 59.85
ATOM	. 2 2 3	N	ASN	159	23.303	25.294	59.502	1.00 40.00
ATOM	:224	CA	ASN	159	23.582	23.944	59.012	1.00 36.67
ATOM	1225	c	ASN	159	23.755	24.002	57.500	1.00 34.11
ATOM	. 226	ŏ	ASN	159	23.223	23.167	56.754	1.00 31.69
ATOM	1226 1227 1228	CB	ASN	159	22.431	22.952	59.367	1.00 46.42
	1227		ASN	159	22.842		59.428	1.00 80.46
MOTA	-220	CG.				21.485		
ATOM	1229	OD1	ASN	159	23.850	21.121	60.054	1.00100.00
ATOM	-230	ND2	ASN	159	22.003	20.620	58.854	1.00 58.09
ATOM	1231	N	GLY	160	24.474	25.044	57.062	1.00 22.34
ATOM	1232	CA	GLY	160	24.686	25.247	55.663	1.00 17.58
MOTA	1233	С	GLY	160	26.055	25.791	55.433	1.00 26.75
MOTA	1234	0	GLY	160	26.960	25.664	56.271	1.00 25.57
MOTA	1235	N	ILE	161	26.200	26.395	54.277	1.00 23.28
ATOM	-235	CA	ILE	161	27.442	26.975	53.909	1.00 16.45
ATOM	:237	С	ILE	161	27.200	28.354	53.395	1.00 15.77
ATOM	1238	0	ILE	161	26.118	28.680	52.962	1.00 15.95
ATOM	1239	CB	ILE	161	28.129	26.117	52.864	1.00 19.27
NOTA	1240 1241	CG1	ILE	161	27.237	26.016	51.619	1.00 18.53
MOTA	241	CG2	ILE	161	28.351	24.735	53.445	1.00 21.96
ATOM	1242	CD1	ILE	161	28.009	25.614	50.350	1.00 14.44
ATOM	243	N	LYS	162	28.226	29.169	53.471	1.00 17.96
ATOM	:244	CA	LYS	162	28.187	30.508	52.948	1.00 14.42
	1245	C	LYS	162	29.216	30.524	51.857	1.00 17.73
ATOM	1245		LYS	162	30.249	29.875	51.857	1.00 17.73
ATOM	1246 1247	0		162				
MOTA	-247	CB	LYS.		28.480	31.540	54.055	1.00 18.15
MOTA	-248	CG	LYS	162	27.221	31.963	54.796	1.00 42.08
MOTA	1249	CD	LYS	162	27.493	32.787	56.039	1.00 70.42
ATOM	1250	N	VAL	163	28.911	31.176	50.759	1.00 13.74
ATOM	1251	CA	VAL	163	29.798	31.201	49.629	1.00 11.95
ATOM	1252	С	VAL	163	29.928	32.610	49.103	1.00 19.30
ATOM	1253	0	VAL	163	28.944	33.318	48.983	1.00 19.84
ATOM	1254	CB	VAL	163	29.249	30.268	48.532	1.00 15.89
ATOM	1255	CG1	VAL	163	30.105	30.277	47.261	1.00 12.29
ATOM	1256	CG2	VAL	163	29.029	23.852	49.077	1.00 15.85
ATOM	1257	N	ASN	:64	31.146	32.999	48.733	1.00 14.03
ATOM		CA	ASN	164	31.382	34.310	48.195	1.00 15.53
ATOM	1259	C	ASN	:64	32.396	34.271	47.050	1.00 20.08
ATOM	260	Ö	ASN	54	33.268	33.386	46.988	1.00 23.49
ATOM	1261	CB	ASN	:54	31.732	35.325	49.308	1.00 20.52
ATOM	1262	CG	ASN	.64	33.196	36 607		1.00 89.21
	1263	001	ASN	154		35.697	49.330	1.00 07.22
ATOM	-203			164	34.020	34.987	-9.929	1.00100.00
ATOM	1264	1102	ASN	-04	33.515	35.831	÷8.700	91.40
ATOM	-265	11	PHE	165	32-244	35.207	46.109	1.00 17.27
ATOM	:266	CA	SHE	165	33.133	25.301	44.953	1.00 10.56

MOTA	1267	С	PHE	165	32.751	36.445	44.071	1.00 15.53
ATOM	1268	0	PHE	165	31.686	37.020	44.251	1.00 15.53
ATOM	1269	СВ	PHE	165	33.207	33.960	44.187	1.00 12.36
		CG		165	31.862	33.486	43.622	1.00 14.35
MOTA	1270		PHE				42.293	1.00 14.55
ATOM	1271	CD1		165	31.510	33.749		
ATOM	1273	CD2	PHE	165	30.978	32.757	44.413	
MOTA		CE1	PHE	165	30.300	33.297	41.759	1.00 22.67
MOTA	1274	CE2	PHE	165	29.774	32.282	43.893	1.00 15.78
MOTA	1275	CZ	PHE	165	29.426	32.572	42.573	1.00 16.20
ATOM	1276	24	LYS	166	33.641	36.799	43.132	1.00 10.79
MOTA	1277	CA	LYS	166	33.417	37.864	42.162	1.00 10.74
ATOM	1278	C	LYS	166	33.603	37.344	40.774	1.00 15.95
ATOM	1279	0	LYS	166	34.602	36.727	40.470	1.00 22.80
ATOM	1280	CB	LYS	166	34.387	39.055	42.249	1.00 16.61
ATOM	1281	CG	LYS	166	34.573	39.688	43.573	1.00 18.11
ATOM	1282	CD	LYS	166	35.540	40.875	43.454	1.00 32.56
ATOM	1283	CE	LYS	166	35.272	41.966	44.476	1.00 48.19
ATOM	1284	ΝZ	LYS	166	34.823	41.435	45.782	1.00 85.81
ATOM	1285	N	ILE	167	32.703	37.704	39.911	1.00 9.75
ATOM	1286	CA	ILE	167	32.768	37.340	38.558	1.00 9.35
ATOM	1287	c	ILE	167	33.203	38.542	37.823	1.00 14.36
ATOM	1288	ō	ILE	167	32.811	39.640	38.170	1.00 16.22
ATOM	1289	СВ	ILE	167	31.379	36.929	38.005	1.00 13.16
ATOM	1290	CGI	ILE	167	30.909	35.624	38.669	1.00 13.02
	1291	CG2		167	31.423	36.786	36.472	1.00 7.91
ATOM			ILE		31.423			
MOTA	1292	CD1	ILE	167	31.773	34.415	38.344	
ATOM	1293	24	ARG	168	34.005	38.299	36.815	1.00 12.19
ATOM	1294	CA	ARG	168	34.500	39.308	35.945	1.00 15.07
MOTA	1295	C	ARG	168	33.948	39.122	34.528	1.00 16.64
ATOM	1296	0	ARG	168	34.278	38.156	33.836	1.00 17.70
ATOM	1297	CB	ARG	168	36.024	39.287	35.944	1.00 16.54
ATOM	1298	CG	ARG	168	36.580	39.632	37.321	1.00 25.54
ATOM	1299	CD	ARG	168	37.894	38.910	37.601	1.00 63.52
ATOM	1300	NE	ARG	168	38.380	38.191	36.416	1.00 73.52
ATOM	1301	CZ	ARG	168	38.764	35.926	36.416	1.00 67.92
ATOM	:302	NHI	ARG	168	38.795	36.192	37.527	1.00 57.44
ATOM	:303	NH2	ARG	168	39.192	36.375	35.271	1.00 59.15
ATOM	1304	14	HIS	169	33.090	40.064	34.098	1.00 14.88
ATOM	1305	CA	HIS	169	32.505	40.025	32.758	1.00 13.24
ATOM	1306	C	HIS	169	33.214	41.001	31.839	1.00 12.64
ATOM	1307	0	HIS	169	33.306	42.203	32.121	1.00 14.99
ATOM	1308	CB	HIS	169	30.970	40.374	32.760	1.00 10.46
ATOM	1309	CG	HIS	169	30.097	39.474	33.573	1.00 6.54
ATOM	1310	:701		169	29.724	38.246	33.111	1.00 12.63
ATOM	1311		HIS	169	29.474	39.695	34.764	1.00 10.21
ATOM	1312	CE1		169	28.892	37.718	34.031	1.00 10.53
ATOM	1313	RE2		169	28.734	38.566	35.063	1.00 11.84
ATOM	1314	N	ASN	170	33.691	40.513	30.737	1.00 10.66
ATOM	1315	CA	ASN	170	34.349	41.358	29.812	1.00 15.87
ATOM	1316	c	ASN	170	33.356	+2.224	29.067	1.00 25.06
ATOM	1317	Ö	ASN	170	32.386	41.701	28.537	1.00 16.60
ATOM	1318	CB	ASN	170	35.110	40.550	28.755	
	:319	CG		170	36.245	39.717		1.00 19.60
ATOM ATOM	1320	001	ASN	170			29.312	1.00 18.70
					36.702	38.752	28.684	1.00 48.29
ATOM	1321	102	ASN	170	36.695	40.073	30.480	1.00 19.13
ATOM	1322	3	ILE	171	33.662	43.527	28.947	1.00 18.75
ATOM	1323	CA	ILE	171	32.848	44.460	28.168	1.00 16.74
ATOM	:324	C	ILE	171	33.459	44.638	26.791	1.00 19.51
ATOM	1325	0	ILE	171	34.643	44.596	25.642	00 21.06
ATOM	1326	CB	ILE	171 171	32.713	45.804	28.842	1.00 20.46
ATOM	1327	CG1	ILE		32.089	45.617	30.193	1.00 19.13 1.00 18.75 1.00 16.74 1.00 19.51 1.00 20.46 1.00 24.79 1.00 19.03
MOTA	1328	CG2	TLE	171	31.852	46.727	27.997	1.00 19.03
ATOM	1329	CDI		171	32,630	46.599	:1.229	
ATOM	1330		GLU	172	32.632	44.818	25.804	
ATOM	1 3 3 1	CA	GLU	172	33.034	44.933	14.420	1.00 17.00
MOTA	1332	2	SLU	172	34,110	43.967	04.147	1.00 17.00
ATOM	1333	0	320	172	34.776	45.298	33.125	1.00 29.20

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ATOM	1334	CB GLU	172	31.813	45.165	23.509	1.00 22.46
ATOM	1335	CG GLU	172	31.122	46.531	23.786	1.00 58.53
ATOM	1336	CD GFA	172	29.871	46.783	22.933	1.00100.00
ATOM	1337	OE1 GLU	172	29.415	45.970	22.156	1.00100.00
ATOM	1338	OE2 GLU	172	29.370	47.982	23.149	1.00100.00
ATOM	1339	N ASP	173	34.277	46.934	25.034	1.00 24.41
ATOM	:340	CA ASP	173	35.292	47.978	24.852	1.00 25.03
ATOM	1341	C ASP	173	36.651	47.624	25.455	1.00 33.40
ATOM	1342	O ASP	173	37.561	48.451	25.518	1.00 30.42
ATOM	1343	CB ASP	273	34.822	49.319	25.401	1.00 23.30
MOTA	1344	CG ASP	173	34.743	49.358	26.912	1.00 32.47
MOTA	1345	OD1 ASP	173	34.406	50.355	27.513	1.00 37.58
MOTA	1346	OD2 ASP	173	34.949	48.196	27.504	1.00 49.22
ATOM	1347	N GLY	174	36.766	46.410	25,956	1.00 23.87
ATOM	1348	CA GLY	174	38.019	45.994	26.537	1.00 21.30
ATOM	1349	C GLY	174	38.012	46.090	28.044	1.00 19.99
MOTA	1350	O GLY	174	38.927	45.585	28.709	1.00 20.45
ATOM	1351	N SER	175	36.972	46.767	28.598	1.00 13.88
ATOM	1352	CA SER	175	36.898	46.931	30.034	1.00 8.70
ATOM	1353	C SER	175	36.296	45.728	30.765	1.00 17.30
ATOM ATOM	1354	O SER	175	36.136 36.288	44.655 48.235	30.175	1.00 18.77
ATOM	1356	OG SER	175 175	36.360	48.235	31.865	1.00 24.79
ATOM	1357	N VAL	176	35.953	45.912	32.051	1.00 13.74
ATOM	1358	CA VAL	176	35.415	44.826	32.864	1.00 16.46
ATOM	1359	C 'AL	176	34.191	45.204	33.703	1.00 22.46
ATOM	1350	O VAL	176	34.159	46.254	34.334	1.00 21.31
ATOM	1361	CB VAL	176	36.477	44.285	33.818	1.00 24.43
ATOM	1362	CG1 VAL	176	35.847	43.344	34.827	1.00 27.45
MOTA	1363	CG2 VAL	176	37.532	43.536	33.035	1.00 25.65
ATOM	1364	N GLN	177	33.234	44.269	33.787	1.00 15.47
ATOM	1365	CA GLN	177	32.048	44.430	34.647	1.00 15.40
ATOM	1366	C GLN	177	32.102	43.457	35.813	1.00 10.60
MOTA	:367	O GLN	177	32.027	42.243	35.634	1.00 13.65
ATOM	1368	CB GLN	177	30.709	44.283	33.872	1.00 15.57
ATOM	1369	CG GLN	177	29.468	44.294	34.828	1.00 19.13
ATOM	:370	CD GLN	177	29.108	45.673	35.361	1.00 14.91
ATOM	1371	OE1 GLN	177	28.759	46.588	34.574	1.00 20.17
ATOM	1372	NE2 GLN	177	29.128	45.821	36.690	1.00 17.28
ATOM	1373	N LEU	178	32.227	43.993	37.018	1.00 8.17
ATOM ATOM	1375	CA LEU	178 178	32.313 30.954	43.180 42.786	38.181	1.00 16.66
ATOM	1376	O LEU	178	30.033	43.608	38.753	1.00 14.66
ATOM	1377	CB LEU	178	33.089	43.896	39.293	1.00 20.63
ATOM	1378	CG LEU	178	34.286	43.110	39.815	1.00 20.63
ATOM	1379	CD1 LEU	178	33.831	42.087	40.852	1.00 45.14
ATOM	1380	CD2 LEU	178	35.018	42.426	38.648	1.00 39.52
ATOM	1381	N ALA	179	30.869	41.550	39.171	1.00 16.72
ATOM	1382	CA ALA	179	29.652	41.033	39.754	1.00 15.55
MOTA	1383	C ALA	179	29.932	40.277	41.040	1.00 15.70
ATOM	:384	O ALA	179	30.337	39.119	41.028	1.00 15.91
ATOM	1385	CB ALA	179	28.853	40.197	38.731	1.00 14.08
ATOM	1386	:: ASP	180	29.694	40.946	42.155	1.00 8.88
ATOM	1387	CA ASP	180	29.897	40.407	43.480	1.00 7.18
ATOM	:388	CASP	180	28.802	39.460	43.891	1.00 17.07
MOTA	1389	O ASP	180	27.651	39-844	43.987	1.00 18.22
ATOM	1390	CB ASP	180	29.934	41.509	44.509	1.00 13.06
ATOM	:391	CG ASP	180	31.285	41.902	44.935	1.00 46.28
ATOM ATOM	1392	OD1 ASP	180	31.981	41.206	45.655	1.00 60.46
ATOM	1393	OD2 ASP	181	31.574	43.121	44.560	1.00 46.61
ATOM	1395	CA HIS	181	29.173 28.213	38.242	44.197	1.00 14.51
ATOM	1396	C HIS	181	28.218	36.897	46.049	
ATOM	1397	O HIS	181	29.255	36.530	46.607	1.00 14.28
ATOM	1398	CB HIS	181	28.450	35.915	43.769	1.00 9.89
ATOM	1399	CG HIS	131	28.077	35.972	42.328	1.00 9.89
ATOM	1400	HD1 HIS	131	28,606	36.926	41.455	1.00 12.24
							12.27

MOTA	1401	CD2	HIS	181	27.279	35.146	41.606	1.00 10.42
ATOM	1402		HIS	181	28.093	26.678	40.269	1.00 9.97
ATOM	1403		HIS	181	27.314	35.594	40.316	1.00 9.38
MOTA	1404	N	TYR	182	27.029	36.897	46.668	1.00 10.40
ATOM	1405	CA	TYR	182	26.848	36.518	48.062	1.00 13.86
ATOM	1406	c	TYR	182	25.871	35.393	48.089	1.00 20.61
ATOM	1407	ō	TYR	182	24.819	35.520	47.532	1.00 16.35
ATOM	1408	CB	TYR	182	26.359	37.664	48.934	1.00 21.12
MOTA	1409	CG	TYR	182	27.421	38.693	49.062	1.00 34.16
ATOM	1410	CD1	TYR	182	27.521	39.715	48.120	1.00 46.06
MOTA	1411	CD2	TYR	182	28.389	38.616	50.064	1.00 16.35 1.00 21.12 1.00 34.16 1.00 46.06 1.00 38.56
MOTA	1412	CE1	TYR	182	28.532	40.674	48.197	1.00 57.53
MOTA	1413	CE2	TYR	182	29.418	39.559	50.147	1.00 40.76
MOTA	1414	cz	TYR	182	29.480	40.594	49.216	1.00 54.61
ATOM	1415	OH	TYR	182	30.461	41.534	49.308	1.00 61.92
ATOM ATOM	1416	N CA	GLN	183	26.246	34.277	48.686 48.583	1.00 17.63
ATOM	1417	CA	GLN	183 183	25.410 25.289	33.104		1.00 16.37
ATOM	1418	0	GLN	183	26.260	32.311	49.863 50.623	1.00 21.39 1.00 19.86 1.00 13.33
ATOM	1420	СВ	GLN	183	25.984	32.219	47.422	1.00 13.33
ATOM	1421	CG	GLN	183	25.651	30.688	47.457	1.00 17.38
ATOM	1422	CD	GLN	183	26.411	29.884	46.389	1.00 17.27
MOTA	1423	OE 1	GLN	183	26.975	30.454	45.456	1.00 17.38 1.00 17.27 1.00 13.80
ATOM	1424	NE2	GLN	183	26.361	28.553	46.473	1.00 13.94
ATOM	1425	N	GLN	184	24.080	31.739	50.055	1.00 13.94
ATOM	1426	CA	GLN	184	23.760	30.829	51.168	1.00 16.55
ATOM	1427	С	GLN	184	23.033	29.582	50.658	1.00 13.60
ATOM	1428	0	GLN	184	22.219	29.640	49.747	1.00 18.01
ATOM	1429	CB	GLN	184	22.949	31.444	52.330	1.00 20.11
ATOM	1430	CG	GLN	184	23.364	32.855	52.768	1.00 74.84
ATOM	1431	CD	GLN	184	22.312	33.517	53.657	1.00100.00
ATOM	1432	OE1	GLN	184	21.159	33.054	53.752	1.00 97.99
ATOM	1433	NE2	GLN	184	22.689	34.625	54.286	1.00100.00
ATOM ATOM	1434	N CA	ASN ASN	185 185	23.418 22.831	23.446	51.207 50.887	1.00 14.75
ATOM	1436	C	ASN	185	22.421	27.155	52.166	1.00 13.86 1.00 16.06 1.00 17.39
ATOM	1437	õ	ASN	185	23.176	25.402	53.172	1.00 17.39
ATOM	1438	CB	ASN	185	23.761	26.212	50.119	1.00 15.20
ATOM	1439	CG	ASN	185	24.110	26,696	48.748	1.00 12.75
ATOM	1440	OD1	ASN	185	24.704	27.758	48.592	1.00 12.75
MOTA	1441	ND2	ASN	185	23.830	25.868	47.763	1.00 17.70
ATOM	1442	N	THR	186	21.227	25.941	52.139	1.00 18.01
MOTA	1443	CA	THR	186	20.707	25.227	53.288	1.00 17.40
ATOM	1444	С	THR	186	19.976	24.010	52.824	1.00 23.63
MOTA	1445	0	THR	186	19.389	23.991	51.730	1.00 24.57
MOTA	1446	CB	THR	186	19.856	26.100	54.206	1.00 28.82
ATOM	1447	061	THR	186	18.874	26.752	53.446	1.00 35.65
ATOM ATOM	1448	CG2	THR	186 187	20.753	27.121	54.903	1.00 28.86
ATOM	1450	CA	PRO	187	19.504	22.951	53.620 53.269	1.00 22.40
ATOM	1451	c	PRO	187	17.988	21.757	53.288	1.00 22.41
ATOM	1452	ŏ	PRO	187	17.390	22.518	54.071	1.00 25.07
ATOM	1453	CB	PRO	197	19.977	20,682	54.337	1.00 19.79
ATOM	1454	CG	PRO	187	20.840	21.449	55.338	1.00 26.98
ATOM	1455	CD	PRO	187	20.786	22.918	54.949	1.00 22.04
MOTA	1456	N '	ILE	188	17.382	20.957	52.453	1.00 18.77
ATOM	1457	CA	ILE	188	15.907	20.855	52.407	1.00 20.12
ATOM	1458	C	ILE	188	15.470	19.766	53.389	1.00 31.58
ATOM	1459	0	ILE	198	14.596	19.966 20.574	54.202	1.00 38.58
ATOM	1460	CB	ILE	:88	15.385	20.574	50.991	1.00 21.52
MOTA	1461	CG1	ILE	198	15.555	21.775	50.102	1.00 16.10
ATOM ATOM	1462	CG2	ILE	188	13.916 15.139	20.141	50.981	1.00 28.85
ATOM	1464	UD1	GLY	139	16.142	21.471	48.660 53.352	1.00 15.31
ATOM	1465	CA.	GLY	:89	15.833	17.531	54.283	1.00 32.94
ATOM	1466	C	GLY	139	16.339	17.817	55.702	1.00 40.20
ATOM	1467	Ö	GLY	139	17.016	17.817	55.967	1.00 35.57
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ATOM	1468	N	ASP	190	16.003	16.928	56.617	1.00 49.41 1.00 55.01 1.00 56.16 1.00 58.30
ATOM	:469	CA	ASP	190	16.392	17.047	58.021	1.00 55.01
ATOM	:470	C	ASP	190	17.556	16.115	58.338	1.00 56.16
ATOM	1471	0	ASP	190	18.083	16.100	59.463	1.00 58.30
ATOM	:472	C8	ASP	190	15.195	16.734	58.955	1.00 63.89
MOTA	1473	CG	ASP	190	14.592	15.365	58.686	1.00 99.€7
ATOM	1474	OD1	ASP	190	14.599	14.466	59.514	1.00100.00
MOTA	1475	OD2	ASP	190	14.088	15.240	57.470	1.00100.00
ATOM	1476	N	GLY	191	17.921	15.312	57.323	1.00 47.20
MOTA	1477	CA	GLY	191	19.015	14.347	57.419	1.00 44.96
MOTA	1478	С	GLY	191	20.359	15.044	57.587	1.00 34.43
ATOM	1479	0	GLY	191	20.452	16.266	57.438	1.00 29.96
ATOM	1480	N	PRO	192	21.402	14.264	57.905	1.00 27.26
ATOM ATOM	1481 14 <b>8</b> 2	CA	PRO	192 192	22.737	14.834	58.100 56.787	1.00 20.55
ATOM	1483	Ö	PRO	192	23.444	14.648	55.740	1.00 23.84
ATOM	1484	CB	PRO	192	23.583	13.764	58.825	1.00 21.00
ATOM	1485	CG	PRO	192	22.739	12.501	58.915	1.00 27.49
ATOM	1486	CD	PRO	192	21.330	12.863	58.448	1.00 27.26
ATOM	1487	N	VAL	193	24.193	16.363	56.892	1.00 17.87
ATOM	1488	CA	VAL	193	24.964	16.902	55.792	1.00 19.51
ATOM	1489	С	VAL	193	26.380	17.108	56.249	1.00 22.37
ATOM	1490	0	VAL	193	26.663	17.189	57.443	1.00 23.84
MOTA	1491	CB	VAL	193	24.449	18.245	55.256	1.00 25.24
ATOM	1492	CG1	VAL	193	23.059	18.118	54.632	1.00 21.90
ATOM	1493	CG2	VAL	193	24.497	19.322	56.346	1.00 24.81
MOTA MOTA	1494	N	LEU	194 194	27.253	17.241	55.277	1.00 19.04
ATOM	1495 1496	CA	LEU	194	28.654 29.006	17.438 18.930	55.516 55.571	1.00 18.71
ATOM	1497	ō	LEU	194	28.907	19.615	54.591	1.00 20.13
ATOM	1498	ČВ	LEU	194	29.412	16.806	54.327	1.00 22.92
ATOM	1499	CG	LEU	194	29.994	15.423	54.542	1.00 30.60
ATOM	1500	CD1	LEU	194	29.227	14.642	55.595	1.00 35.19
ATOM	1501	CD2	LEU	194	30.048	14.672	53.211	1.00 25.61
ATOM	1502	N	LEU	195	29.453	19.430	56.713	1.00 17.39
ATOM	1503	CA	LEU	195	29.881	20.808	56.785	1.00 18.83
ATOM	1504	C	LEU	195	31.389	20.837	56.579	1.00 28.32
ATOM	1505	0	LEU	195	32.161	20.152	57.281	1.00 21.98
ATOM	1506	CB	LEU	195	29.489	21.525	58.072	1.00 22.20
ATOM ATOM	1507 1508	CG CD1	LEU	195 195	28.055 27.937	21.349 21.508	58.444 59.941	1.00 26.40
ATOM	1509	CD2	LEU	195	27.225	22.395	57.726	1.00 26.90
ATOM	1510	N N	PRO	196	31.789	21.610	55.597	1.00 21.58
ATOM	1511	CA	PRO	196	33.177	21.666	55.154	1.00 22.17
ATOM	1512	c	PRO	196	34.080	22.623	55.892	1.00 29.56
ATOM	1513	0	PRO	196	33.635	23.588	56.490	1.00 29.04
ATOM	1514	CB	PRO	196	33.054	22.265	53.752	1.00 22.77
ATOM	1515	CG	PRO	196	31.761	23.104	53.735	1.00 18.99
ATOM	1516	CD	PRO	196	30.910	22.567	54.861	1.00 16.42
ATOM	1517	N	ASP	197	35.379	22.410	55.716	1.00 22.95
ATOM	1518	CA	ASP	197 197	36.364 36.556	23.370	56.134 54.931	1.00 19.71
ATOM	1520	o	ASP	197	36,251	23.913	53.800	
ATOM	1521	CB	ASP	197	37.711	22.730	56.446	1.00 24.88
ATOM	1522	CG	ASP	197	37.690	21.913	57.687	1.00 43.93
ATOM	1523	OD1	ASP	197	36.912	22.117	58.608	1.00 53.47
ATOM	1524	OD2	ASP	197	38,634	21.006	57.694	1.00 31.58
ATOM	1525	M	ASN	198	37.062	25.501	55.168	1.00 31.58
ATOM	1526	CA	ASN	198	27.254	26.470	54.118	1.00 15.38
ATOM	1527	c	ASN	198	37.974	25.889	52.971	1.00 19.61
ATOM	1528	0	ASN	198	38.958	25.236	53.134	1.00 22.69
ATOM	1529	CB	ASN	198	39.012	27.704	54.614	1.00 24.48
ATOM ATOM	1530 1531	CG OD1	ASN ASN	198 198	37.236	28.504	55.632	1.00 52.21
ATOM	1532	ND2		198	17.854	29.556	55.961 55.150	1.00 55.11
MOM	:533	11	HIS	199	27.462	25.125	11.801	1.00 16.30
MOTA	1534	CA.	HIS	199	33.071	25.627	30.616	1.00 15.30
	.554						.5.516	12.50

ATOM	1535	c	HIS	199	37.496	26.357	49.450	1.00 14.85
ATOM	1536	0	HIS	199	36.757	27.295	49.643	1.00 16.45
ATOM	1537	CB	HIS	199	37.988	24.103	50.471	1.00 16.53
ATOM	1538	CG	HIS	199	36.597	23.628	50.218	1.00 16.65
ATOM	1539	::01		199	35.695	23.491	51.244	1.00 17.85
ATOM	1540	CD2		199	35.987	23.282	49.048	1.00 18.67
ATOM	1541	CEI	HIS	199	34.561	23.052	50.688	1.00 19.45
ATOM	1542	::E2	HIS	199	34.716	22.905	49.364	1.00 18.74
ATOM	1543	24	TYR	200	37.879	25.998	48.247	1.00 12.56
MOTA	1544	CA	TYR	200	37.334	26.689	47.100	1.00 14.01
MOTA	1545	C	TYR	200	37.207	25.824	45.870	1.00 15.57
ATOM	1546	0	TYR	200	37.793	24.751	45.768	1.00 20.20
ATOM	1547	CB	TYR	200	38.030	28.011	46.779	1.00 19.79
ATOM	1548	CG	TYR	200	39.382	27.745	46.202	1.00 22.25
ATOM	1549	CD1	TYR	200	39.543	27.526	44.835	1.00 22.53
ATOM	1550	CD2	TYR	200	40.473	27.605	47.057	1.00 25.73
ATOM	1551	CEI	TYR	200 200	40.800	27.222	44.317	1.00 35.51
ATOM ATOM	1552 1553	CE2	TYR	200	41.739 41.896	27.314	46.559	1.00 29.34
ATOM	1554	OH	TYR	200	43.153	26.820	45.186 44.703	1.00 62.66
ATOM	1555	y	LEU	201	36.393	26.309	44.703	1.00 15.07
ATOM	1556	ĊA	LEU	201	36.147	25.680	43.678	1.00 11.01
ATOM	1557	CA	LEU	201	36.753	26.532	42.593	1.00 17.30
ATOM	1558	ŏ	LEU	201	36.619	27.753	42.610	1.00 20.19
ATOM	1559	ĊВ	LEU	201	34.628	25.518	43.354	1.00 10.09
ATOM	1560	CG	LEU	201	33.749	25.027	44.480	1.00 13.41
ATOM	1561	CD1		201	32.293	24.938	43.954	1.00 17.11
ATOM	1562	CD2	LEU	201	34.196	23.635	44.927	1.00 23.03
ATOM	1563	31	SER	202	37.407	25.868	41.651	1.00 10.75
ATOM	1564	CA	SER	202	38.047	26.490	40.528	1.00 8.51
ATOM	1565	С	SER	202	37.222	26.189	39.294	1.00 11.56
ATOM	1566	0	SER	202	36.919	25.038	38,996	1.00 14.58
ATOM	1567	CB	SER	202	39.485	25.987	40.442	1.00 15.68
ATOM	1568	OG	SER	202	40.067	26.353	39.228	1.00 36.44
MOTA	1569	:1	THR	203	36.798	27.241	38,601	1.00 12.36
ATOM	1570	CA	THR	203	35.879	27.067	37.499	1.00 15.60
MOTA	1571	С	THE	203	36.417	27.521	36.195	1.00 20.19
ATOM	1572	0	THR	203	37.192	28.472	36.114	1.00 18.29
ATOM	1573	CB	THR	203	34.565	27.892	37.757	1.00 20.51
ATOM	1574	OG1	THR	203	34.911	29.260	37.780	1.00 20.39
ATOM	1575	CG2	THR	203	33.935	27.557	39.093	1.00 6.80
MOTA	1576	::	GLN	204	35.913	26.883	35.164	1.00 10.30
MOTA MOTA	1577 1578	CA	GLN	204 204	36.173 34.956	27.271	33.807	1.00 14.85
ATOM	1579	Ö	GLN	204	34.334	26.980 25.932	32.921	1.00 23.14
ATOM	1580	C3	GLN	204	37.475	26.696	33.056	1.00 21.66
ATOM	1581	CG	GLN	204	37.271	25.371	33.237 32.518	1.00 20.33
MOTA	1582	CD	GLN	204	38.588	24.722	32.518	1.00 40.16
ATOM	1583	OE1	GLN	204	39.011	24.716	31.035	1.00 41.80
ATOM	1584	HE2	GLN	204	39.276	24.241	33.235	1.00 34.80
ATOM	1585	31	SER	205	34.619	27.913	32.021	1.00 15.83
ATOM	:586	CA	SER	205	33.447	27.762	31.172	1.00 14.60
ATOM	1587	C	SER	205	33.654	28.307	29.783	1.00 20.21
ATOM	1588	0	SER	205	34.282	29.337	29.581	1.00 17.82
ATOM	1589	C3	SER	205	32.197	28.445	31.758	1.00 11.88
ATOM	1590	OG	'SER	205	32.121	28.406	33.177	1.00 15.45
MOTA	1591	::	ALA	206	33.065	27.630	28.827	1.00 13.00
MOTA	1592	SA	ALA	206	33.079	23.029	27.426	1.00 9.99
ATOM	1593	Ξ.	ALA	206	31.623	23.192	26.924	1.00 21.23
ATOM	1594	0	ALA	206	30.809	27.306	27.139	1.00 14.10
ATOM	1595	23	ALA	206	33.751	26.936	26.596	1.00 13.45
ATOM	1596	11	LEU	207	31.335	39.320	26.263	1.00 16.09
ATOM	1597	CA	LEU	207	30.036	29.617	25.706	1.00 12.07
MOTA	1598	5	LEU	207	30.070	29.445 29.840	24.235	1.00 19.76
ATOM	1500	.0	LEU	207	31.014	2.840	23.576	1.00 20.82
ATOM ATOM	1601	-2	150	207	29.580	11.057 31.493	26.004	1.00 8.24
ATUM	- 401	- 3	LEU	207	29.744	37	27.457	1.00 16.38

FIG 5-25

ATOM : 668 CA ASP

32.790 ..00 1602 207 28.955 27.707 13.78 ATOM CD1 LEU 1603 ..00 30.406 28.400 18.79 ATOM CD2 LEU 207 29.268 ATOM 1604 N SER 208 29.011 28.863 23.698 1.00 15.35 208 22.270 1.00 13.74 ATOM 1605 CA SER 28.914 28.692 21.794 ATOM 1606 С SER 208 27.449 28.852 1.00 20.16 22.594 1607 208 26.548 29.085 1.00 A TYOM 0 SER 15.81 ATON :608 C8 SER 208 29.495 27.367 21.822 1.00 17.82 ATOM 1609 SER 208 28.769 26.311 22.431 1.00 31.45 OG ATOM 1610 LYS 209 27.242 28.738 20.485 1.00 16.50 ATOM 209 25.907 28.828 19.906 1.00 18.02 1611 CA LYS 1.00 29.99 ATON 1612 С LYS 209 25.637 27.610 19.031 ATOM 1613 0 LYS 209 26.578 27.004 18,502 1.00 32.55 ATOM 1614 CB LYS 209 25.783 30.100 19.082 1.00 20.96 CG 209 24.746 31.055 1.00 34.50 ATON 1615 LYS 19.606 ATOM 1616 CD LYS 209 25.262 31.964 20.666 1.00 22.72 33.159 20.896 1.00 18.96 ATOM 1617 CF TYS 209 24,370 ATOM 1618 N7 LYS 209 23.565 33.067 22.116 1.00 27.39 ATOM 1619 N ASP 210 24.347 27.241 18.912 1.00 27.01 ATOM 1620 CA ASP 23.890 26.159 18.038 1.00 24.62 23.465 ATOM 1621 ASP 210 26.793 16.705 1.00 26.77 22.468 ATOM 1622 n ASP 210 27.514 16.605 1.00 23.00 ATOM 1623 CB ASP 210 25.361 18.691 1.00 24.43 ATOM 1624 CG ASP 210 22.197 24.249 17.839 1.00 35.53 ATOM 1625 OD1 ASP 210 22.333 24.185 16.631 1.00 36.53 ATOM 1626 OD2 210 23.400 1.00 45.51 ASF 21.499 18.535 1.00 30.25 ATOM 1627 N PRO 211 24.306 26.618 15.708 1.00 30.30 ATOM 1628 CA PRO 211 24.120 27.224 14.397 ATOM 1629 C PRO 211 22.733 26.982 13.770 1.00 39.72 1.00 ATOM 1630 0 PRO 211 22.253 27.782 12.959 37.65 1.00 29.99 ATOM 1631 CB PRO 211 25.197 26.620 13.500 ATOM 1632 ĊĞ PRO 211 25.782 25.418 14.255 1.00 38.59 ATOM 1633 CD PRO 211 25.158 25.405 15.647 1.00 35.05 ATOM 1634 ASN 22.102 25.868 39.64 М 212 14.140 1.00 112 212 ATOM :635 CA ASN 20.808 25.515 13.592 1.00 39.60 ATOM 1636 С ASN 19.642 25.894 14.497 1.00 41.92 ATOM 1637 0 ASN 212 212 212 18.485 25.518 14.263 1.00 42.30 24.028 ATOM 1638 CB ASN 20.789 13.236 1.00 48.64 1.00 53.61 ATOM 1639 CG ASN 21.883 23.678 12.230 ATOM 1640 N GLU 19.947 26.675 1.00 27.84 213 15.520 ATOM 1641 CB 18.953 16.478 1.00 20.43 ATOM 1642 GLU 213 18.485 28.527 16.241 1.00 29.95 ATOM 1643 a GLU 19.247 29.475 16.324 1.00 32.77 ATOM 1644 CB GLU 213 19.535 26.878 17.894 1.00 16.45 ATOM 1645 CG GLU 213 18.594 27.326 18.995 1.00 18.29 ATOM 1646 CD GLU 213 17.229 26.703 18.853 1.00 38.01 ATOM 1647 OE1 CIT 213 16.238 27.334 18.508 1.00 25.07 OE2 1.00 ATOM 1648 GLU 213 17.223 25.423 19.122 19.17 ATOM 1649 N LYS 214 17.223 28.713 15.963 1.00 22.99 1650 CA 1.00 ATOM LYS 214 16.721 30.081 15.726 22.84 ATOM 1651 c 1.75 214 214 214 16.252 30.778 16.982 1.00 21.50 ō 32.016 ATOM 1652 L.YS 16.130 17.032 28.15 1.00 1653 CB LYS 15.653 ATOM 30.197 14.606 1.00 27.58 ATOM 1654 CC 16.153 LYS 214 29.816 13.209 1.00 32.71 2 16.752 ATOM 1655 LYS 214 30.979 12.431 1.00 55.31 ATOM 1656 21 ARG 115 15.947 30.028 13.014 1.00 14.52 ATOM 1657 Ch ARG 315 15.518 30.726 19.209 1.00 15.58 ATOM 1658 С ARG :15 16.719 31.382 19.892 ..00 21.87 17.848 ATOM 1659 0 ARG 215 31.075 19.572 1.00 26.69 ATOM 1660 CB 250 14.808 23.804 20.159 .00 18.82 ATOM 1661 CG ARG 13.660 29.067 19.475 1.00 23.30 CD ATON 1662 ARG 115 13.220 27.806 19.205 ..00 15.45 ARG 14.107 ATOM 1663 UF 215 26.668 1.00 28.08 CZ ARG 115 ATOM 1664 14.022 25.473 20.543 ..00 21.38 115 ATOM :665 SH 1 ARG 13.074 28.215 11.455 .00 23.92 ATOM :666 WH2 ARG 14.893 24.514 20.223 1.00 20.46 16.466 ATOM 1667 \*1 ASP 10.830 ..00 16.72

17.556

12.895

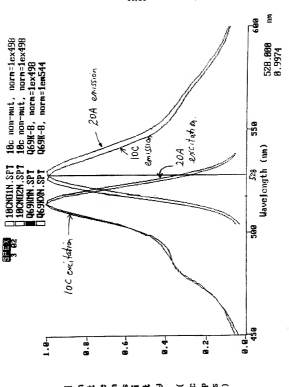
19.06

ATOM	1669	С	ASP	216	18.047	31.817	22.607	1.00 20.02
ATOM	1670	ŏ	ASP	216	17.261	31.214	23.350	1.00 18.45
ATOM	1671	CB	ASP	216	17.066	34.169	22.383	:.00 21.33
ATOM	1672	CG	ASP	216	18.138	35.140	22.893	1.00 20.97
ATOM	1673	001		216	17.869	36.079	23.620	1.00 28.46
ATOM	1674	OD2		216	19.342	34.900	22.441	1.00 20.37
ATOM	1675	N	HIS	217	19.332	31.537	22.589	1.00 13.18
ATOM	1676	CA	HIS	217	19.813	30.482	23.433	1.00 11.21
ATOM	1677	c	HIS	217	21.313	30.614	23.723	1.00 21.35
ATOM	1678	ō	HIS	217	22.014	31.471	23,163	1.00 15.03
ATOM	1679	CB	HIS	217	19.587	29.168	22.690	1.00 13.03
ATOM	1680	CG	HIS	217	20.525	29.025	21.542	1.00 15.49
ATOM	1681		HIS	217	20.463	29.871	20,449	1.00 17.88
ATOM	1682		HIS	217	21.589	28.172	21.361	1.00 17.51
ATOM	1683	CE1	HIS	217	21.457	29.524	19.635	1.00 17.94
ATOM	1684	NE2	HIS	217	22.152	28.501	20.151	1.00 17.59
ATOM	1685	N	MSE	218	21.794	29.725	24.576	1.00 11.26
ATOM	1686	CA	MSE	218	23.186	29.642	24.887	1.00 11.49
ATOM	1687	С	MSE	218	23.560	28.198	25.094	1.00 24.15
ATOM	1688	0	MSE	218	22.822	27.446	25.751	1.00 20.70
MOTA	1689	CB	MSE	218	23.539	30.421	26.172	1.00 12.84
MOTA	1690	CG	MSE	218	24.809	30.004	26.907	1.00 12.59
ATOM	1691	SE	MSE	218	25.267	31.128	28.434	1.00 29.94
MOTA	1692	CE	MSE	218	24.039	30.502	29.781	1.00 13.54
ATOM	1693	N	VAL	219	24.727	27.824	24.558	1.00 15.62
ATOM	1694	CA	VAL	219	25.309	26.518	24.782	1.00 10.58
MOTA	1695	С	VAL	219	26.473	26.689	25.753	1.00 16.54
ATOM	1696	0	VAL	219	27.280	27.604	25.585	1.00 15.54
ATOM	1697	CB	VAL	219	25-774	25.883	23.498	1.00 15.08
ATOM	1698	CG1	VAL	219	26.330	24.495	23.824	1.00 14.34
ATOM	1699	CG2	VAL	219	24.599	25.766	22.512	1.00 15.78
ATOM	1700	N	LEU	220	26.523	25.836	26.783	1.00 10.95
ATOM	1701	CA	LEU	220	27.490	25.939	27.850	1.00 11.01
ATOM	1702	С	LEU	220	28.206	24.643	28.184	1.00 21.26
ATOM	1703	0	LEU	220	27.592	23.577	28.324	1.00 15.94
ATOM	1704	CB	LEU	220 220	26.807 27.624	26.545 26.578	29.100 30.402	1.00 13.75
MOTA	1706	CD1		220	28.433	27.875	30.483	1.00 21.10
ATOM ATOM		CD2	LEU	220		26.556	21.586	1.00 22.04
ATOM	1707 1708	N CD2	LEU	221	26.663 29.570	24.758	28.273	1.00 19.04
ATOM	1709	CA	LEU	221	30.498	23.666	28.697	1.00 13.22
MOTA	1710	c	LEU	221	31.309	24.178	29.887	1.00 10.73
ATOM	1711	ō	LEU	221	31.846	25.267	29.857	1.00 12.98
ATOM	1712	CB	LEU	221	31.382	23.102	27.549	1.00 13.74
ATOM	1713	CG	LEU	221	32.580	22.257	28.045	1.00 18.64
ATOM	1714	CD1		221	32.149	20.868	28.496	1.00 17.38
ATOM	1715	CD2		221	33.571	22,109	26.911	1.00 26.97
ATOM	1716	N	GLU	222	31.316	23.446	30.963	1.00 9.31
ATOM	1717	CA	GLU	222	31.936	23.929	32.144	1.00 9.97
ATOM	1718	c	GLU	222	32.548	22.803	32.951	1.00 12.94
ATOM	1719	0	GLU	222	32.072	21.662	32.966	1.00 13.38
MOTA '	1720	CB	GLU	222	30.836	24.762	32.896	1.00 12.14
ATOM	1721	CG	GLU	222	31.092	25.119	34.364	1.00 13.88
ATOM	1722	CD	GLU	222	29.895	25.891	34.934	1.00 13.57
MOTA	1723		GLU	222	29.128	26.477	34.240	1.00 19.47
ATOM	1724		GLU	222	29.752	25.789	36.207	1.00 18.51
MOTA	1725	N	PHE	223	33.687	23.123	33.542	1.00 15.86
ATOM	1726	CA	PHE	223	34.476	22.227	34.373	1.00 9.34
ATOM	1727	c	PHE	223 223	34.711	22.864	35.722	1.00 11.08
ATOM	1728	0	PHE	223	35.028	24.055	35.828	1.00 19.86 1.00 3.30 1.00 10.50
ATOM	1729	CB	PHE	223	35.847	21.919	33.684	1.00 8.30
ATOM	1730	CG	PHE	223	35.703	21.134	32,431	00 10.50
ATOM	1731	CD1		223	35.570	19.747	32.469	1.00 13.56
ATOM	1732	CD2		123 123 123	35.750	21.750	31.184	
ATOM	1733			23	35.481	19.010	21.287	1.00 12.58
ATOM	1734			123	35.667	21.032	19.995	
ATOM	1735	CZ	PHE	123	35.521	19.648	30.050	1.00 10.27

ATOM	1736	21	VAL	224	34.542	22.081	36.765	1.00 9.23
ATOM	1737	CA	VAL	224	34.708	22.587	38.080	1.00 11.13
ATOM	1738	С	VAL	224	35.324	21.553	39.010	1.00 17.52
ATOM	:739	0	VAL	224	34.848	20.418	39.137	1.00 13.17
MOTA	1740	CB	VAL	224	33.370	23.078	38.662	1.00 16.51
ATOM	1741	CG1	VAL	224	33.622	23.736	40.022	1.00 13.90
ATOM	1742	CG2	VAL	224	32.674	24.048	37.697	1.00 13.85
ATOM	1743	N	THR	225	36.380	21.965	39.676	1.00 11.71
ATOM	:744	CA	THR	225	37.026	21.099	40.517	1.00 11.51
ATOM	1745	С	THR	225	37.366	21.798	41.927	1.00 14.76
ATOM	1746	0	THR	225	37.702	23.002	41.962	1.00 16.64
ATOM	1747	CB	THR	225	38.162	20.279	40.014	1.00 20.38
ATOM	1748	OG1	THR	225	39.288	20.337	40.822	1.00 30.44
ATOM	1749	CG2	THR	225	38.468	20.722	38.631	1.00 10.89
ATOM	1750	И	ALA	226	37.222	21.065	43.011	1.00 7.89
ATOM	1751	CA	ALA	226	37.478	21.595	44.352	1.00 11.53
MOTA	1752	С	ALA	226	38.969	21.558	44.677	1.00 16.61
ATOM	1753	0	ALA	226	39.687	20.699	44.199	1.00 15.60
MOTA	1754	CB	ALA	226	36.695	20.847	45.444	1.00 12.17
ATOM	1755	N	ALA	227	39.395	22.490	45.479	1.00 13.95
ATOM	1756	CA	ALA	227	40.789	22.550	45.871	1.00 19.64
ATOM	1757	С	ALA	227	40.987	23.299	47.170	1.00 26.23
MOTA	1758	0		227	40.042		47.840	
MOTA	1759	CB	ALA	227	41.557	23.246	44.760	1.00 18.42 1.00 23.29
ATOM	1760 1761	N CA	GLY	228 228	42.245 42.616	24.292	47.523 48.658	1.00 23.29
ATOM	1762	CA	GLY	228	42.805	23.562	49.939	1.00 32.93
ATOM	1763	ŏ	GLY	228	42.948	24.201	51.009	1.00 32.93
ATOM	1764	N	ILE	229	42.803	22.231	49.842	1.00 33.59
ATOM	1765	CA	ILE	229	43.006	21.375	50.998	1.00 31.81
ATOM	1766	c	ILE	229	44.016	20.291	50.633	1.00 28.78
ATOM	1767	ŏ	ILE	229	45.090	20.176	51.246	1.00 96.02
ATOM	:768	C3	EEE	229	41.691	20.772	51.519	1.00 35.70
ATOM	1769	CG1	ILE	229	40.890	21.807	52.325	1.00 30.66
ATOM	1770	CG2	ILE	229	41.990	19.549	52.392	1.00 33.37
ATOM	1771	CD1	ILE	229	39.386	21.715	52.092	1.00 38.74
ATOM	1772	0	HOH	301	27.530	12.735	38,010	1.00 15.09
ATOM	:773	0	HOH	302	23.919	34.589	37.331	1.00 10.29
ATOM	1774	0	HOH	303	27.229	34.816	35.487	1.00 11.12
ATOM	1775	0	HOH	304	29.914	18.943	44.692	1.00 15.10
ATOM	1776	0	HOH	305	30.956	21.886	49.900	1.00 21.47
ATOM	1777	0	HOH	306	20.072	31.196	43.592	1.00 15.85
ATOM	1778	0	HOH	307	26.660	48.630	33.797	1.00 24.67
ATOM	1779	0	HOH	308	22.329	33.239	41.399	1.00 14.11
ATOM	1780	0	нон	309	22.465	48.025	32.810	1.00 18.51
ATOM	1781	0	нон	310	31.012	39.126	29.118	1.00 15.01
ATOM	1782	0	HOH	311	33.067	35.809	33.010	1.00 19.92
ATOM	1783	0	HOH	312	31.130	37.076	30.841	1.00 12.58
ATOM	1784	0	НОН	313 314	40.304 34.166	30.058	38.616	1.00 56.07
ATOM	1785 1786	0	HOH	315	36.215	26.379	57.222	1.00 22.58
MOTA	1787	0	HOH	316	33.866	35.320 29.786	43.598	1.00 22.30
ATOM	1865	0	нон	317	42.341	20.166		1.00 12.21
ATOM	1788	0	HOH	318	10.270	28.684	43.534	1.00 26.67
ATOM	1789	ŏ	HOH	319	28.448	16.822	30.403	1.00 43.65
ATOM	790	0	нон	320	30.612	20.922	37.231	1.00 21.57
ATOM	:791	ŏ	HOH	321	11.639	37.421	26.801	1.00 21.57
ATOM	792	ŏ	нон	322	27.030	37.308	36.869	1.00 13.10
ATOM	1793	ö	HOH	323	33.119	14.524	43.070	1.00 30.93
ATOM	794	ō	HOH	324	37.973	14.036	53.352	1.00 35.39
ATOM	1795	ō	HOH	325	32.015	49.100	37.028	1.00 59.37
ATOM	:796	0	HOH	326	11.959	12.020	43.429	1.00 29.06
ATOM	1797	0	HOH	327	36.760	29.941	31.666	1.00 22.03
ATOM	:864	0	HOH	328	15.305	26.513	15.694	1.00 39.62
MOTA	1798	0	HOH	329	33.005	46.924	36.994	1 00 22 27
MOTA	1363	0	HOH	330	13.501	36.134	22.715	1.00 45.20
MOTA	:799	0	HOH	331	33.609	31.296	16.261	1.00 23.65

ATOM	1862	0	нон	332	34.942	24.780	29.532	1.00 38.93
ATOH	1800	ō	HOH	333	25.235	12.919	54.611	1.00 36.20
ATOM	1861	ō	HOH	334	38.048	23.467	36.645	1.00 37.73
ATOM	1801	ŏ	нон	335	12.284	43.511	38.338	1.00 33.79
ATOM	1802	ŏ	HOH	336	9.826	47.020	32.568	1.00 46.67
ATOM	1803	ō	HOH	337	7.671	41.532	29.806	1.00 40.88
ATOM	1804	0	HOH	338	15.430	23.713	26.808	1.00 34.73
ATOM	1805	ŏ	HOH	339	24.344	20.385	25.121	1.00 53.42
ATOM	1806	ŏ	HOH	340	31.550	10.656	40.819	1.00 47.85
ATOM	1807	0	HOH	341	17.569	23.030	25.796	1.00 28.17
ATOM	1808	ŏ	HOH	342	19.174	38.552	23.965	1.00 45.54
ATOM	1809	ŏ	HOH	343	24.268	37.527	25.415	1.00 30.97
ATOM	1810	ŏ	HOH	344	21.266	29.482	41.551	1.00 19.69
ATOM	1811	ō	HOH	345	20.668	26.999	41.933	1.00 11.81
ATOM	1812	ŏ	HOH	346	24.780	24.795	43.460	1.00 20.95
ATOM	1813	ŏ	HOH	347	42.962	13.170	46.312	1.00 31.00
ATOM	1814	ō	нон	348	32.322	14.088	47.013	1.00 28.20
ATOM	1815	ō	HOH	349	31.708	13.186	49.679	1.00 35.57
ATOM	1816	ŏ	HOH	350	22.408	35.801	50.514	1.00 40.71
ATOM	1817	ŏ	HOH	351	25.366	47.090	42.583	1.00 38.15
ATOM	1818	ŏ	HOH	352	27.243	47.647	43.977	1.00 41.55
ATOM	1819	o	HOH	353	29.868	45.076	42.906	1.00 29.32
ATOM	1820	ŏ	HOH	354	14.175	22.269	42.680	1.00 74.11
ATOM	1821	Ö	HOH	355	13.414	10.739	35.791	1.00 29.92
ATOM	1822	o	HOH	356	20.338	9.974	37.765	1.00 30.46
ATOM	1823	ő	HOH	357	23.520	40.420	24.953	1.00 29.75
ATOM	1824	ŏ	HOH	358	25.718	41.692	26.023	1.00 30.43
ATOM	1825	ŏ	HOH	359	26.826	38.466	25.345	1.00 31.72
ATOM	1826	ŏ	HOH	360	37.768	42.373	25.123	1.00 41.53
ATOM	1827	ō	HOH	361	40.078	42.268	25.852	1.00 37.12
ATOM	1828	ŏ	нон	362	31.483	38.677	22.083	1.00 54.21
ATOM	1829	ŏ	HOH	363	33.891	37.723	30.126	1.00 23.35
ATOM	1860	ŏ	HOH	364	39.936	26.543	36.329	1.00 47.93
ATOM	1830	ō	HOH	365	36.631	34.210	41.636	1.00 62.74
ATOM	1831	ő	HOH	366	37.038	29.783	52.197	1.00 40.07
ATOM	1832	ŏ	HOH	367	37.289	37.407	40.231	1.00 37.59
ATOM	1833	ŏ	HOH	368	18.930	17.517	52.472	1.00 35.80
ATOM	1834	ŏ	HOH	369	19.506	18.914	57.913	1.00 45.72
ATOM	1835	ŏ	HOH	370	30.903	25.708	41.139	1.00 21.54
ATOM	1836	ŏ	нон	371	30.369	25.678	24.583	1.00 22.46
ATOM	1837	ō	нон	372	21.000	33.705	20.826	1.00 26.00
ATOM	1838	ō	HOH	373	13.648	32.794	21.329	1.00 27.98
ATOM	1839	ō	нон	374	29.735	25.683	38.707	1.00 21.00
ATOM	1859	ō	нон	375	33.670	24.419	60.503	1.00 50.04
ATOM	1840	o	HOH	376	30.034	11.047	37.420	1.00 43.28
ATOM	1841	0	HOH	377	8.662	35.846	35.068	1.00 51.94
ATOM	1842	ā	нон	378	10.847	36.466	39.503	1.00 42.32
ATOM	1843	Ó	HOH	379	14.395	48.943	39.085	1.00 29.72
ATOM	1844	0	HOH	380	36.676	11.660	40.172	1.00 39.81
MOTA	1845	0	HOH	381	35.968	7.212	34.763	1.00 58.66
ATOM	1846	0	нон	382	17.426	21.988	21.077	1.00 41.69
-ATOM	1847	0	HOH	383	29.837	22.623	39.378	1.00 32.82
ATOM	1848	0	нон	384	23.855	29.386	55.164	1.00 55.00
ATOM	1849	0	HOH	385	17.408	35.360	47.495	1.00 61.61
ATOM	1850	0	HOH	386	27,900	49.720	42.448	1.00 47.70
ATOM	1851	0	HOH	387	13.932	36.230	44.385	1.00 45.08
ATOM	1852	0	HOH	388	12.650	23.021	43.288	1.00 49.86
ATOM	1853	0	HOH	389	16.974	42.367	43.435	1.00 34.38
ATOM	1854	0	HOH	390	37.335	42.653	28.295	1.00 64.46
ATOM	1855	0	HOH	391	29.701	49.856	35.323	1.00 62.61
ATOM	1856	0	HOH	392	27.267	50.835	33.976	1.00 66.60
MOTA	1857	0	HOH	393	19.661	29.181	51.537	1.00 34.01
ATOM	1858	0	HOH	394	29.412	17,505	59.089	1.00 51.78
TER								
END								





Normalized

International application No. PCT/US97/14593

A. CLASSIFICATION OF SUBJECT MATTER  IPC(6) : CO713 21/04; CO7K 14/00, 16/00; CI2N 1/20, 15/00, 15/09, 15/63  US CL : Please See Extra Sheet.  According to International Patent Classification (IPC) or to both national classification and IPC	
According to international Patent Classification (IPC) or to both national classification and IPC	
TO PERSON OF THE	
B. FIELDS SEARCHED  Minimum documentation searched (classification system followed by classification symbols)	
U.S. : 435/252.3, 252.33, 325, 410, 320.1; 530/350, 387; 536/23.1, 23.4	
Documentation searched other than minimum documentation to the extent that such documents are included in the field	ds searched
Electronic data base consulted during the international search (name of data base and, where practicable, search to Please See Extra Sheet.	rms used)
C. DOCUMENTS CONSIDERED TO BE RELEVANT	
Category* Citation of document, with indication, where appropriate, of the relevant passages Relevant	at to claim No.
X WO 96/23810 A1 (THE REGENTS OF THE UNIVERSITY OF CALIFORNIA) 08 August 1996, abstract and claims.	
1-41,	54-7
X HEIM et al. Improved green fluorescence. Nature. Vol. 373, 23 88 February 1995, pages 663-664, see Figure on page 664.	10-16, 18-
26, 28 39-41	3-32, 34-37,
	•
X Further documents are listed in the continuation of Box C. Soe patent family annex.	
* Special easegories of cited documents: "I" inter document yeldshede after the international fall.  *A* document defining the general state of the ort which is not considered to be of particular relevance or the orthogonal fall.  **Description of the orthogonal fall of the orthogonal fall of the orthogonal fall of the principle or theory underlying the internation.	g date or priority and to understand
*8" earlier document published on or after the intermetional filling date  "X" document of perticular relevance; the claimed involve  "L" document which may throw doubts on priority obtain(s) or which is  "B" document of perticular relevance; the claimed involve  "the document is the document in taken alone	ention cannot be an inventive stop
1." dooesen't which may throw doubt on priority chain(s) or which is tied to madelick the profession that or mother desirance other "O" desirance referring to an ord dischours, use, achibition or other manufacture to provide the provided of the provid	he document is
"P" document published prior to the international filing data but later than "A." document member of the same patent family the priority data claimed	
Date of the actual completion of the international search 18 DECEMBER 1997  Date of mailing of the international search report 2 7 JAN 1998	
Name and mailing address of the ISA/US Commissioner of Patents and Trademarks  Authorized officer  Twfor	
Box PCT         NASHAAT T. NASHED           Washington, D.C. 20231         NASHAAT T. NASHED           Facsimile No. (703) 305-3230         Telephone No. (703) 308-0196	

International application No. PCT/US97/14593

Category*	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No
Category	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No
X  Y	HEIM et al. Wavelength mutations and posttranslational autoxidation of green fluorecent protein, Proc. Natl. Acad. Sci. USA. Vol. 91, December 1994, pages 12501-12504, see abstract.	46, 48, 50, 52, 54, 56, 94 
Y	PEROZZO et al. X-ray diffraction and time-resolved fluorescence analysis of Aequorea green fluorescent crystals. Journal of Biological Chemistry. 05 June 1988, Vol. 263, No. 16, pages 7713-7716.	1-41, 46-57, 86- 94
X  Y	DELAGRAVE et al. Red-shifted excitation mutants of the green fluorescent protein. Bio/Technology. February 1995, Vol. 13, pages 151-153, see Table 1 on page 152.	46-57, 86-88, 91  1-41
Y	EHRIG et al. Green-fluorescent protein mutants with altered fluorescence excitation spectra. FEBS Letters. 1995, Vol. 367, pages 163-166, abstract.	1-41, 46-57, 89, 90
Y	WANG et al. Implication for bcd mRNA localization from spatial distribution of exu protein in Drosophila oogenesis. Nature. 02 June 1994, Vol. 369, 400-403, see Figure 1.	32-41, 54-57
Р, Ү	ORMO et al. Crystal structure of the Aquorea victoria green fluorescent protein. Science. 06 September 1996, Vol. 273, pages 1392-1395, abstract.	1-41, 46-57, 86- 94
P, Y	YANG et al. The molecular structure of green fluorescent protein. Nature Biotechnology. October 1996, Vol. 14, pages 1246-1251, abstract.	1-41, 46-57, 86- 94
P, Y	PALM et al. The structural basis for spectral variations in green fluorescent protein. Nature Struct. Biol. May 1997, Vol. 4, Number 5, pages 361-365.	1-41, 46-57, 86- 94
A	US 5,491,084 A (CHALFIE et al.) 13 February 1996, entire document.	1-41, 46-57, 86- 94

International application No. PCT/US97/14593

Box 1 Ob	servations where certain claims were found unsearchable (Continuation of Item 1 of first sheet)
This interna	tional report has not been established in respect of certain claims under Article 17(2Xa) for the following reasons:
	claims Nos.: ecause they relate to subject matter not required to be scarched by this Authority, namely:
	<u>♥</u>
<u>ا</u> ا	Claims Nos.:  coasse they relate to parts of the international application that do not comply with the prescribed requirements to such a extent that no meaningful international search can be carried out, specifically:
	Claims Nos.: ecause they are dependent claims and are not drafted in accordance with the accord and third sentences of Rule 6.4(a).
Box 11 Of	pervations where unity of invention is lacking (Continuation of Item 2 of first sheet)
This Intern	ational Searching Authority found multiple inventions in this international application, as follows:
Pleas	se See Extra Sheet.
	•
	s all required additional search fees were timely paid by the applicant, this international search report covers all searchable
	is an required admindral search tees were unterly part by the appricant, this international search report covers an searchance laims.
	s all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment f any additional fee.
ه ت	s only some of the required additional search fees were timely paid by the applicant, this international search report covers sly those claims for which fees were paid, specifically claims Nos.: 4.6-57, 8.6-94
,	
	o required additional search fees were timely paid by the applicant. Consequently, this international search report is stricted to the invention first mentioned in the claims; it is covered by claims Nos.:
	·
Remark on	Protest X The additional search fees were accompanied by the applicant's protest.  No protest accompanied the payment of additional search fees.

International application No. PCT/US97/14593

A. CLASSIFICATION OF SUBJECT MATTER:

US CL :

435/252.3, 252.33, 325, 410, 320.1; 530/350, 387; 536/23.1, 23.4

B. FIELDS SEARCHED

Electronic data bases consulted (Name of data base and where practicable terms used):

APS, STN: Medline, Caplus, Sciscarch, Lifesci, Biosis, Embase, Wpids, Biotechds.

Search terms: acquorea and green fluorescent, T-203, Thr-203, T203, DNA, cDNA, sequence, s651, 1203h, s651, 1203y, s72a, f641, s65g, 203y, s72a, s65g, v681, t203y, 142x, v61x, t62x, v68x, q69x,n121x,y145x, v150x, f165x, i167x, q183x

BOX II. OBSERVATIONS WHERE UNITY OF INVENTION WAS LACKING

This ISA found multiple inventions as follows:

This application contains the following inventions or groups of inventions which are not so linked as to form a single inventive concept under PCT Rule 13.1. In order for all inventions to be searched, the appropriate additional search fees must be paid.

Group I, claims 1-41, drawn to DNA coding for mutant fluorescent green protein having mutation at Thr-203, the fluorescent protein anatibody labeled with the fluorescent protein and the DNA coding for a fusion protein consisting of an antibody and the mutant fluorescent protein.

Group II, claims 42-45 and 58-61, drawn to a DNA probe labled with mutants fluorescent.

Group III, claims 46-57 and 86-94, drawn to drawn to DNA coding for mutants fluorescent green protein having mutation at an amino acid residue other than Thr-203, the fluorescent protein, antibody labeled with the fluorescent protein and the DNA coding for a fusion protein consisting of an antibody and the mutant fluorescent protein.

Group IV, claims 62-64 and 68-70, drawn to a method for engineering fluorescent protein.

Group V, claims 65-67, drawn to method of producing fluorescent resonance energy transfer.

Group VI, claims 71-74, drawn to a fluorescent protein crystal baving the amino acid sequence SEQ ID NO: 2.

Group VII, claims 75-82, drawn to a computation method for the design of fluorescent protein.

Group VIII, claims 83-85, drawn to a storage device containing the atomic coordinate.

Group IX, claims 95-100, drawn to a method of identifying test chemicals.

The inventions listed as Groups I-IX do not relate to a single inventive concept under PCT Rule 13.1 because, under PCT Rule 13.2, they lack the same or corresponding special technical features for the following reasons:

Each of the above group has a special technical feature defined by the first claim in the Group. The following are the special technical feature for each Group: (a) Group 1 is the succlose acid coding for fluorescent protein having at least the mutation at Thr-203, (b) Group II is a fluorescent DNA probe labeled with a mutant fluorescent protein, (c)Group III is the nucleic soil sequence coding for mutant fluorescent protein having mutation at residues other than T-203, (d) Group IV is a method for the engineering of mutant fluorescent proteins, (e) Group V is a method for producing fluorescent resonance energy transfer, (d) Group V is the protein crystal of the widd-type protein, (e) Group VIII is the computation method to design mutant fluorescent protein with different fluorescent characteristics, (h) Group VIII is a storage device for data, and (i) Group IV is a method of identifying test chemically

Group I encompasses the nucleic soid coding for the mutant fluorescent protein, expression vector, recombinant bost cell, the mutant proteins and a use for the DNA in making the fluiton protein consisting of antibody and the fluorescent protein. Group II represent a second use for the mutant protein of Group I. Also, the special technical feature of Group I is different from that of Group III because the DNA of each Group codes for different sets of mutants that do not share common feature. The special technical feature of this Group I is different sets of Groups FUIX.

The special technical feature of Group II, the fluorescent DNA probe is clearly different from those of Groups III-IX.

The DNA probe of Group II represent a second use of the fluorescent protein of Group II.

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The method of engineering fluorescent protein of Group IV is different from that of producing fluorescent resonance energy transfer of Group V because the resulting fluorescence is different in each case and vary in its characteristic Similarly, the special technical features of each of Groups IV and V are different from those of the crystal of Group VI, the computation method of Group VII, the storage device of Group VIII, and the method of identifying chemicals of Group IX. Finally, the crystal of Group VI, the computation method of Group VII, the storage device of Group VIII, and the method of identifying chemicals of Group IX are clearly unrelated to each other and there is no special technical feature that connects them together.